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(54) Title: SOLUTION AND CRYSTAL STRUCTURES OF MMP-13 ACTIVE SITE AND USES THEREOF

(57) Abstract: The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), as well as to (i) methods of using the MMP-13 structure to rationally design or identify compounds or molecules that inhibit or activate MMP-13 activity, and (ii) compounds identified using said methods.

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SOLUTION AND CRYSTAL STRUCTURES OF MMP-13 ACTIVE SITE AND USES THEREOF

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Field of the Invention

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), as well as to (i) methods of using the MMP-13 structure to rationally design or identify compounds or molecules that inhibit or activate MMP-13 activity, and (ii) compounds identified using said methods.

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Background of the Invention

Human collagenase-3 (MMP-13) is a member of the matrix metalloproteinase (MMP) family which includes the collagenases, stromelysins and gelatinases. The MMPs are involved in the degradation of the extracellular matrix and are associated with normal tissue remodeling processes such as pregnancy, wound healing, and angiogenesis. MMP expression and activity is highly controlled because of the degradative nature of these enzymes, where an apparent loss in MMP regulation results in the pathological destruction of connective tissue and the ensuing disease state. Accordingly, MMPs are a highly active set of targets for the design of therapeutic agents for the disease areas of arthritis and oncology (for reviews, see Woessner, J. F., FASEB 1991; Ries, C., and Petrides, E., Biol. Chem. Hoppe-Seyler 1995; Browner, M. F., Perspect. Drug Discovery Des. 1995; Morphy, et al., Curr. Med. Chem. 1995; and Zask, et al., Curr. Pharm. Des. 1996).

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MMP-13 was identified on the basis of differential expression in normal breast tissues and in breast carcinoma. In addition, its expression has been reported in squamous cell carcinomas of the larynx, head and neck, in HCS-2/8 human chondrosarcoma cells, during fetal ossification, and in articular cartilage of arthritic patients.

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There have been a number of X-ray and NMR structures solved for the catalytic domain of MMPs complexed with a variety of inhibitors (see e.g., Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Science 1994; Lovejoy, et al., Ann. N. Y. Acad. Sci. 1994; Lovejoy, et al.,

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Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl. Acad. Sci. U.S.A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. USA 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr. 1996; Gooley, et al., J. Biomol. NMR 1996; Betz, et al., Eur. J. Biochem. 1997; Gonnella, et al., Bioorg. Med. Chem. 1997; and Moy, et al., Biochemistry 1998). There is a close similarity in the overall threedimensional fold for these proteins consistent with the relatively high sequence homology (> 40%). Despite this similarity in the MMP structures, there is a distinct substrate specificity between these enzymes indicative of specific 10 biological roles for the various MMPs and a corresponding association with unique disease processes. One example of this potential specificity is the overexpression of MMP-13 in breast carcinoma and MMP-1 in papillary carcinomas. Therefore, the current paradigm in the development of MMP inhibitors is to design specificity into the structures of the small molecule instead of developing 15 a broad spectrum MMP inhibitor (Birkedal-Hansen, et al., Crit. Rev. Oral Biol. Med. 1993; and Rockwell, et al., J. Am. Chem. Soc. 1996). The rationale behind this approach is that an inhibitor specific for the MMP uniquely associated with a disease process may potentially minimize toxic side effects. Therefore, extensive structural information for the various MMPs is critical for a 20 structure-based approach in designing inhibitor selectivity (Birkedal-Hansen, et al., Crit. Rev. Oral Biol. Med. 1993; Rockwell, et al., J. Am. Chem. Soc. 1996; Ghose, et al., J. Am. Chem. Soc. 1995; Hajduk, et al., J. Am. Chem. Soc. 1997; and Olejniczak, et al., J. Am. Chem. Soc. 1997).

This concept has been facilitated by the extensive structural data available for the MMPs where a significant difference in the size and shape of the S1' pocket has been observed (Moy, et al., Biochemistry 1998; Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Ann. N.Y. Acad. Sci. 1994; Lovejoy, et al., Biochemistry 1994; Lovejoy, et al., Science 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl.

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Acad. Sci. U.S.A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. U.S.A. 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr. 1996; Gooley, et al., J. Biomol. NMR 1996; Betz, et al., Eur. J. Biochem. 1997; and Gonnella, et al., Bioorg. Med. Chem. 1997). This structural difference across the MMP family provides an obvious approach for designing specificity into potent MMP inhibitors by designing compounds that appropriately fill the available space in the S1' pocket while taking advantage of sequence differences. A number of examples have been previously reported using this approach where some selectivity between MMPs has been achieved by incorporating a biphenyl into the S1' pocket (see e.g., Hajduk, et al., J. Am. Chem. Soc. 1997; and Olejniczak, et al., J. Am. Chem. Soc. 1997).

The inventors have determined both the solution and crystal structures of MMP-13, and, using rational drug design methods, have designed a novel, potent inhibitor that is highly selective for MMP-13.

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Summary of the Invention

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), and more specifically, to the crystal and solution structures of MMP-13 complexed with the inhibitor N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide (hereinafter referred to as "Compound A"), as determined using crystallography, spectroscopy and various computer modeling techniques. Particularly, the invention is directed to an MMP-13 active site comprised of the three dimensional structures of various binding pockets located both to the right (S1', S2', S3') and left (S1, S2, S3) of the catalytic zinc of MMP-13, and most particularly is directed to the three dimensional structure of an MMP-13 active site comprising the catalytic zinc and the S1' binding pocket, which is critical to the design and selection of inhibitors with increased potency and specificity for MMP-13, or conversely, for the design and selection of inhibitors of matrix metalloproteinases that are specific against MMP-13.

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Accordingly, the present invention discloses a solution comprising a biologically active catalytic fragment of human collagenase-3 (MMP-13) complexed with Compound A, as well as a crystallized catalytic fragment of MMP-13 complexed with Compound A. The three dimensional structure of the catalytic fragment of MMP-13 is provided by the relative atomic structural coordinates of Figure 4, as obtained from spectroscopy data, and Figure 5, as obtained from crystallography data. Also provided is an active site of MMP-13, characterized by a catalytic zinc, a beta strand, a Ca²⁺ binding loop, an alpha helix and a random coil region, wherein the beta strand of said active site preferably comprises residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1, the Ca²⁺ binding loop comprises residues F75, D76, G77, P78, and S79 according to Figure 1, the alpha helix comprises residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1, and the random coil region comprises residues P139, I140, and Y141 according to Figure 1. Said active site is further characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, ± a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å.

In an alternate embodiment of the invention, an active site of MMP-13 is characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å.

The solution or crystal structural coordinates of MMP-13 or portions thereof as provided by this invention may be stored in a

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machine-readable form on a machine-readable storage medium, e.g. a computer hard drive, diskette, DAT tape, etc., for display as a three-dimensional shape or for other uses involving computer-assisted manipulation of, or computation based on, the structural coordinates or the three-dimensional structures they define. By way of example, the data defining the three dimensional structure of MMP-13 or an MMP-13 complex of the present invention, or of a portion of MMP-13 or an MMP-13 complex as disclosed herein, may be stored in a machine-readable storage medium, and may be displayed as a graphical three-dimensional representation of the relevant structural coordinates, typically using a computer capable of reading the data from said storage medium and programmed with instructions for creating the representation from such data.

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Accordingly, the present invention provides a machine, such as a computer, programmed in memory with the coordinates of the MMP-13 molecule or molecular complex, or portions thereof (such as, by way of example, the coordinates of the MMP-13 catalytic zinc with adjacent S1', S2' and/or S3' binding pockets), together with a program capable of converting the coordinates into a three dimensional graphical representation of the structural coordinates on a display connected to the machine. A machine having a memory containing such data aids in the rational design or selection of inhibitors or activators of MMP-13 activity, including the evaluation of ability of a particular chemical entity to favorably associate with MMP-13 or an MMP-13 complex as disclosed herein, as well as in the modeling of compounds, proteins, complexes, etc. related by structural or sequence homology to MMP-13.

The present invention is additionally directed to a method of determining the three dimensional structure of a molecule or molecular complex whose structure is unknown, comprising the steps of first obtaining crystals or a solution of the molecule or molecular complex whose structure is unknown, and then generating X-ray diffraction data from the crystallized molecule or molecular complex and/or generating NMR data from the solution of the molecule or molecular complex. The generated diffraction or spectroscopy data from the molecule or molecular complex can then be compared with the known

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three dimensional structure of MMP-13 as disclosed herein, and the three dimensional structure of the unknown molecule or molecular complex conformed to the known MMP-13 structure using standard techniques such as molecular replacement analysis, 2D, 3D and 4D isotope filtering, editing and triple resonance NMR techniques, and computer homology modeling. Alternatively, a three dimensional model of the unknown molecule may be generated by generating a sequence alignment between MMP-13 and the unknown molecule, based on any or all of amino acid sequence identity, secondary structure elements or tertiary folds, and then generating by computer modeling a three dimensional structure for the molecule using the three dimensional structure of, and sequence alignment with, MMP-13.

The present invention further provides a method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of using a three dimensional structure of MMP-13 as defined by the relative structural coordinates of amino acids encoding MMP-13 to design or select a potential inhibitor or activator, and synthesizing or obtaining said potential inhibitor or activator. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of an empty MMP-13 active site in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors. The method of the present invention is preferably used to design or select inhibitors of MMP-13 activity.

Alternatively, the present invention provides a method for identifying a potential inhibitor or activator that is selective for one or more members of the matrix metalloproteinase family except MMP-13, comprising the steps of (i) using the three dimensional structures of MMP-13 and the desired target matrix metalloproteinase(s) as defined by the relative structural coordinates of amino acids encoding MMP-13 and the target matrix metalloproteinase(s) in order to design or select such a potential inhibitor or

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activator, and (ii) synthesizing or obtaining said potential inhibitor or activator. In this case, the potential inhibitor or activator is designed to incorporate chemical or steric features favorable for association with an active site of the desired matrix metalloproteinase(s) and unfavorable for association with an MMP-13 active site, preferably where said active site comprises the MMP-13 S1' pocket. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of empty MMP-13/matrix metalloproteinase active sites in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors.

Also provided by the present invention are the inhibitors and activators designed or selected using the methods disclosed herein.

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Brief Description of the Figures

Figure 1 depicts the amino acid sequence encoding the catalytic fragment of human MMP-13.

Figure 2 depicts the sequence based alignment between (A) MMP-13 and MMP-8 and (B) MMP-13 and MMP-1 used for the MMP-13 homology model.

Figure 3 is an illustration of the sulfonamide derivative of the hydroxamic inhibitor N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide (Compound A), with the corresponding proton labels.

Figure 4 lists the atomic structure coordinates for the restrained minimized mean structure of MMP-13 complexed with Compound A as derived by NMR spectroscopy. "Atom type" refers to the atom whose coordinates are being measured. "Residue" refers to the type of residue of which each measured atom is a part - i.e., amino acid, cofactor, ligand or solvent. The "x, y and z" coordinates indicate the Cartesian coordinates of each measured atom's location

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(Å). All non-protein atoms (Compound A, zinc and calcium) are listed as HETATM instead of atoms using PDB conventions.

Figure 5 lists the atomic structure coordinates for MMP-13 as derived by X-ray diffraction of a crystallized MMP-13:Compound A complex. Figure headings are as noted above, except "Occ" indicates the occupancy factor, and "B" indicates the "B-value", which is a measure of how mobile the atom is in the atomic structure (Ų). "MOL" indicates the segment identification used to uniquely identify each molecule in the crystal.

Figure 6 is an illustration of the Compound B inhibitor, with the corresponding proton labels.

Figure 7 is a design scheme dividing 2-[Benzyl-(4-phenethyloxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide (hereinafter referred to as "Compound C") into two components corresponding to its potency component (2-[Benzyl-(4-methoxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, hereinafter referred to as "Compound D") and its selectivity component, thereby providing the basis for the design of a hybrid inhibitor with Compound B.

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Figure 8A is a design scheme showing the flow from Compound B and Compound C to the hybrid inhibitor benzofuran-2-carboxylic acid (2-{4-20 [benzyl-(2-hydroxycarbamoyl-4,6-dimethyl-phenyl)-sulfamoyl]-phenoxy}-ethyl)-amide (hereinafter referred to as "Compound E"). Figure 8B illustrates an expanded view of the NMR MMP-13:Compound B complex overlayed with the MMP-13:Compound D model, demonstrating the approach to forming the hybrid inhibitor Compound E. The MMP-13 active site is shown as a grid surface with Compound B and Compound D shown as liquorice bonds. The view is looking at the S1' pocket.

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Detailed Description of the Invention

As used herein, the following terms and phrases shall have the meanings set forth below:

"Compound A" is N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)pyridin-3-ylmethyl-amino]-3-methyl-benzamide, as shown in Figure 3.

"Compound B" is the compound having the chemical structure shown in Figure 6. "Compound C" is 2-[Benzyl-(4-phenethyloxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, as shown in Figure 7. "Compound D" is 2-[Benzyl-(4-methoxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, also shown in Figure 7. "Compound E" is Benzofuran-2-carboxylic acid (2-{4-[benzyl-(2-hydroxycarbamoyl-4,6-dimethyl-phenyl)-sulfamoyl]-phenoxy}-ethyl)-amide, as shown in Figure 8A. "Compound F" is 2-(Benzyl-4-(3-phenyl-propoxy)-benzenesulfonyl]-amino)-N-hydroxy-3,5-dimethyl-benzamide.

Unless otherwise noted, "MMP-13" includes both human collagenase 3 as encoded by the amino acid sequence of Figure 1 (including conservative substitutions thereof), as well as "MMP-13 analogues", defined herein as proteins comprising an MMP-13 like active site as defined by the present invention, including, but not limited to, an active site characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, ± a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å. Alternatively, an MMP-13 analogue of the present invention is a protein which comprises an MMP-13 like active site characterized by a catalytic zinc, a beta strand, a Ca²⁺ binding loop, an alpha helix and a random coil region, or, more particularly, comprising an active site characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and of amino acid residues N14, L15, T16, Y17, R18, I19, V20,

F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, or more preferably, where said three dimensional structure further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to Figures 4 or 5, in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms (N, C α , C, and O) of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

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Unless otherwise indicated, "protein" or "molecule" shall include a protein, protein domain, polypeptide or peptide.

"Structural coordinates" are the Cartesian coordinates corresponding to an atom's spatial relationship to other atoms in a molecule or molecular complex. Structural coordinates may be obtained using x-ray crystallography techniques or NMR techniques, or may be derived using molecular replacement analysis or homology modeling. Various software programs allow for the graphical representation of a set of structural coordinates to obtain a three dimensional representation of a molecule or molecular complex. The structural coordinates of the present invention may be modified from the original sets provided in Figures 4 or 5 by mathematical manipulation, such as by inversion or integer additions or subtractions. As such, it is recognized that the structural coordinates of the present invention are relative, and are in no way specifically limited by the actual x, y, z coordinates of Figures 4 and 5. Further, it is recognized that the structural coordinates taken from Figure 5 may be from either molecule of MMP-13 catalytic fragment in the MMP-13:Compound A crystal (i.e., from A-13 or B-13).

An "agent" shall include a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug.

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"Root mean square deviation" is the square root of the arithmetic mean of the squares of the deviations from the mean, and is a way of expressing deviation or variation from the structural coordinates described herein.

It will be obvious to the skilled practitioner that the numbering of the amino acid residues in the various isoforms of MMP-13 or in MMP-13 analogues covered by the present invention may be different than that set forth herein, or may contain certain conservative amino acid substitutions that yield the same three dimensional structures as those defined by Figures 4 or 5 herein. Corresponding amino acids and conservative substitutions in other isoforms or analogues are easily identified by visual inspection of the relevant amino acid sequences or by using commercially available homology software programs. "Conservative substitutions" are those amino acid substitutions which are functionally equivalent to the substituted amino acid residue, either by way of having similar polarity, steric arrangement, or by belonging to the same class as the substituted residue (e.g., hydrophobic, acidic or basic), and includes substitutions having an inconsequential effect on the three dimensional structure of MMP-13 with respect to the use of said structure for the identification and design of MMP-13 activators or inhibitors, for molecular replacement analyses and/or for homology modeling.

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An "active site" refers to a region of a molecule or molecular complex that, as a result of its shape and charge potential, favorably interacts or associates with another agent (including, without limitation, a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug). As such, the active site may include both the actual site of substrate cleavage or collagenase activity, as well as certain or all binding sites or pockets adjacent to the site of substrate cleavage that nonetheless may affect MMP-13 activity upon interaction or association with an agent, either by direct interference with the site of substrate cleavage or by indirectly affecting the steric conformation or charge potential of the MMP-13 molecule. The catalytic center of the MMP-13 molecule is characterized by a zinc atom chelated by H119, H123 and H129. MMP-13 binding sites or pockets located to the right of

the catalytic zinc include S1', S2' and S3'. Binding sites or pockets to the left of the catalytic zinc include S1, S2 and S3.

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13) or an MMP-13 analogue, and more specifically, to the crystal and solution structures of MMP-13 complexed with an inhibitor, referred to herein as "Compound A", as determined using crystallography, spectroscopy and various computer modeling techniques. The three dimensional solution and crystal structures of the MMP-13:Compound A complex (as disclosed herein at Figures 4 or 5, respectively) and the uncomplexed MMP-13 catalytic fragment (which may be computationally derived from the structural coordinates of Figures 4 or 5) are useful for a number of applications, including, but not limited to, the visualization, identification and characterization of MMP-13 active sites, including the MMP-13 catalytic zinc chelated by H119, H123 and H129, as well as the various MMP-13 binding pockets adjacent to the catalytic zinc of the MMP-13 molecule. The active site structures may then be used to predict the orientation and binding affinity of a designed or selected activator or inhibitor of the MMP-13 protein. Accordingly, the invention is particularly directed to the three dimensional structure of an MMP-13 active site, including but not limited to the S1', S2', S3', S1, S2 and/or S3 binding pockets, taken separately or together with the catalytic zinc of the MMP-13 molecule.

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The present invention provides a solution comprising a biologically active catalytic fragment of human collagenase-3 (MMP-13) complexed with Compound A. In a particular embodiment, the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1, or conservative substitutions thereof. Preferably, the solution provided for herein comprises MMP-13 complexed with Compound A in a 1:1 molar ratio, and more preferably comprises 1 mM MMP-13 in an equimolar complex with Compound A, in a buffer comprising 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl₂, 0.1mM ZnCl₂, 2mM NaN₃, and 10 mM deuterated DTT in either 90% H₂O/10% D₂O or 100% D₂O, at a preferred pH of 6.5. The concentration of

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MMP-13:Compound A in the solution should be high enough to yield a good signal-to-noise ratio in the NMR spectrum, but not so high as to result in precipitation or aggregation of the protein. Further, the MMP-13 of the solution may be either ¹⁵N enriched or ¹⁵N, ¹³C enriched. As exemplified below, NMR spectra from the solution of the present invention are preferably obtained at a temperature of 35°C.

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The secondary structure of the catalytic fragment used in the solution of the present invention comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands, configured in the order β_{I} , α_{A} , β_{II} , β_{III} , β_{IV} , β_{V} , α_{B} , and α_{C} . The three alpha helices correspond to residues 28-44 (α_{A}), 112-123 (α_{B}) and 153-163 (α_{C}) of Figure 1, and the five beta strands correspond to residues 83-86 (β_{I}), 95-100 (β_{II}), 59-66 (β_{III}), 14-20 (β_{IV}), and 49-53 (β_{V}) of Figure 1, respectively. While the solution of the present invention comprises MMP-13 in a 1:1 molar ratio with Compound A, it is understood that one of ordinary skill in the art may devise additional solutions using alternate inhibitors or ligands in the appropriate molar concentrations, thereby preventing the auto-degradation of MMP-13 and creating a solution of sufficient stability and concentration to obtain a usable NMR spectrum.

MMP-13, as well as MMP-13 analogues, where said protein comprises an active site characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the solution coordinates of Figure 4, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å. These residues comprise the residues most closely associated with Compound A in the MMP-13:Compound A complex, as determined from the observed NOEs between MMP-13 and Compound A (Table

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Alternatively, a protein used in the solution of the present invention comprises an active site characterized by a catalytic zinc, a beta strand (comprising amino acid residues N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca²⁺ binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, characterized by a three dimensional structure comprising the relative structural 10 coordinates of the catalytic zinc and the amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figure 4, or more preferably, where said three dimensional structure further comprises the relative structural coordinates of amino acid residues G80, L81, 15 L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figure 4 (incorporating an S1' pocket in the active site), or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to 20 Figure 4 (further defining a hydrophobic area at the bottom of the S1' pocket), including, in each case, conservative substitutions of said amino acids and, in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms (N, Ca, C, and O) of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). Finally, in the most preferred embodiment, the protein used in the solution of the present invention comprises the complete structural coordinates according to Figure 4, \pm a root mean square deviation from the conserved backbone atoms of said amino acids (or conservative substitutions thereof) of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å).

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Also provided by the present invention is a crystallized catalytic fragment of MMP-13 complexed with Compound A. The crystal of the present invention effectively diffracts X-rays for the determination of the structural coordinates of the MMP-13:Compound A complex, and is characterized as being in orthorhombic form with space group P21212, and having unit cell parameters of a=108.3Å, b=79.8Å, and c=36.1Å. Further, the crystal complex of the present invention consists of two molecules of MMP-13:Compound A complex in the asymmetric crystal unit.

In a preferred embodiment, the MMP-13 of the crystal complex of the present invention comprises the amino acid residues of Figure 1 (or conservative substitutions thereof), and is characterized by a secondary structure comprising three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands, configured in the order β_I , α_A , β_{II} , β_{III} , β_{IV} , β_V , α_B , and α_C . Further, the three alpha helices preferably correspond to residues 28-44 (α_A), 112-123 (α_B) and 153-163 (α_C) of Figure 1, and the five beta strands correspond to residues 83-86 (β_I), 95-100 (β_{II}), 59-66 (β_{III}), 14-20 (β_{IV}), and 49-53 (β_V) of Figure 1, respectively.

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The protein used in the crystal or crystal complex of the present invention includes MMP-13, as well as MMP-13 analogues, where said protein comprises an active site characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the crystal coordinates of Figure 5, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å.

Alternatively, a protein used in the crystal or crystal complex of the present invention comprises an active site characterized by a catalytic zinc, a beta strand (comprising amino acid residues N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca²⁺ binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions

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thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figure 5, or more preferably, where said three dimensional structure further comprises the relative structural coordinates of amino acid 10 residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figure 5 (incorporating an S1' pocket in the active site), or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to Figure 5 (further defining a hydrophobic area at the bottom of the S1' pocket), in each case, including conservative substitutions of the said amino acids and, in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 20 0.5Å).

Finally, in the most preferred embodiment, the protein used in the crystal of the present invention comprises the complete structural coordinates according to Figure 5, \pm a root mean square deviation from the conserved backbone atoms of said amino acids (or conservative substitutions thereof) of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å).

Molecular modeling methods known in the art may be used to identify an active site or binding pocket of the MMP-13 molecule, MMP-13 molecular complex, or an MMP-13 analogue. Specifically, the structural coordinates provided by the present invention may be used to characterize a

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three dimensional model of the MMP-13 molecule, molecular complex or MMP-13 analogue. From such a model, putative active sites may be computationally visualized, identified and characterized based on the surface structure of the molecule, surface charge, steric arrangement, the presence of reactive amino acids, regions of hydrophobicity or hydrophilicity, etc. Such putative active sites may be further refined using chemical shift perturbations of spectra generated from various and distinct MMP-13 complexes, competitive and non-competitive inhibition experiments, and/or by the generation and characterization of MMP-13 mutants to identify critical residues or characteristics of the active site.

The identification of putative active sites of a molecule or molecular complex is of great importance, as most often the biological activity of a molecule or molecular complex results from the interaction between an agent and one or more active sites of the molecule or molecular complex. Accordingly, the active sites of a molecule or molecular complex are the best targets to use in the design or selection of activators or inhibitors that affect the activity of the molecule or molecular complex.

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The present invention is directed to an active site of MMP-13 or an MMP-13 analogue, that, as a result of its shape, reactivity, charge potential, etc., favorably interacts or associates with another agent (including, without limitation, a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug). As such, the active site of the present invention includes both the actual site of substrate cleavage or collagenase activity (the catalytic zinc chelated by H119, H123, and H129), as well as binding sites or pockets adjacent to the site of substrate cleavage (i.e., S1', S2', S3', S1, S2, and/or S3) that may nonetheless affect MMP-13 activity upon interaction or association with an agent, either by direct interference with the site of substrate cleavage or by indirectly affecting the steric conformation or charge potential of the MMP-13 molecule. Accordingly, the present invention is directed to an active site of the MMP-13 molecule characterized by a zinc atom chelated by H119, H123 and H129, and preferably the S1' binding pocket to the right of the catalytic zinc.

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In an alternate embodiment, the active site of the present invention is characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å.

Alternatively, the active site of the present invention is characterized by a catalytic zinc, a beta strand (comprising amino acid residues 10 N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca²⁺ binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, is characterized by a three dimensional structure comprising the relative solution or crystal structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, respectively, or more preferably, where said three dimensional structure further comprises the relative solution or crystal structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, where said three dimensional structure still further comprises the relative solution or crystal structural coordinates of F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å

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(or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

In order to use the structural coordinates generated for a crystal or solution structure of the present invention as set forth in Figures 4 and 5, respectively, it is often necessary to display the relevant coordinates as, or convert them to, a three dimensional shape or graphical representation, or to otherwise manipulate them. For example, a three dimensional representation of the structural coordinates is often used in rational drug design, molecular replacement analysis, homology modeling, and mutation analysis. This is typically accomplished using any of a wide variety of commercially available 10 software programs capable of generating three dimensional graphical representations of molecules or portions thereof from a set of structural coordinates. Examples of said commercially available software programs include, without limitation, the following: GRID (Oxford University, Oxford, 15 UK); MCSS (Molecular Simulations, San Diego, CA); AUTODOCK (Scripps Research Institute, La Jolla, CA); DOCK (University of California, San Francisco, CA); Flo99 (Thistlesoft, Morris Township, NJ); Ludi (Molecular Simulations, San Diego, CA); QUANTA (Molecular Simulations, San Diego, CA); Insight (Molecular Simulations, San Diego, CA); SYBYL (TRIPOS, Inc., St. Louis. MO); and LEAPFROG (TRIPOS, Inc., St. Louis, MO). 20

For storage, transfer and use with such programs, a machine, such as a computer, is provided for that produces a three dimensional representation of the MMP-13 molecule, a portion thereof (such as an active site or a binding site), a MMP-13 molecular complex, or an MMP-13 analogue. The machine of the present invention comprises a machine-readable data storage medium comprising a data storage material encoded with machine-readable data. Machine-readable storage media comprising data storage material include conventional computer hard drives, floppy disks, DAT tape, CD-ROM, and other magnetic, magneto-optical, optical, floptical and other media which may be adapted for use with a computer. The machine of the present invention also comprises a working memory for storing instructions for processing the

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machine-readable data, as well as a central processing unit (CPU) coupled to the working memory and to the machine-readable data storage medium for the purpose of processing the machine-readable data into the desired three dimensional representation. Finally, the machine of the present invention further comprises a display connected to the CPU so that the three dimensional representation may be visualized by the user. Accordingly, when used with a machine programmed with instructions for using said data, e.g., a computer loaded with one or more programs of the sort identified above, the machine provided for herein is capable of displaying a graphical three-dimensional representation of any of the molecules or molecular complexes, or portions of molecules of molecular complexes, described herein.

In one embodiment of the invention, the machine-readable data comprises the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to Figures 4 or 5, in each case, including conservative substitutions thereof, and in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å), wherein said structural coordinates characterize an active site of MMP-13 or an MMP-13 analogue.

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In an alternate preferred embodiment, the machine-readable data comprises the structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, in each case, including conservative substitutions thereof, and in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å). In an even more preferred embodiment, the machine-readable data further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126,

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L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, still further comprises the relative structural coordinates of F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

Finally, it is most preferred that the machine-readable data comprise the relative structural coordinates of all residues constituting the MMP-13 catalytic fragment according to Figures 4 or 5, in each case, \pm a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å. In each case, the noted embodiments comprise conservative substitutions of the noted residues resulting in same structural coordinates within the stated root mean square deviation.

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The structural coordinates of the present invention permit the use of various molecular design and analysis techniques in order to (i) solve the three dimensional structures of related molecules, molecular complexes or MMP-13 analogues, and (ii) to design, select, and synthesize chemical agents capable of favorably associating or interacting with an active site of an MMP-13 molecule or MMP-13 analogue, wherein said chemical agents potentially act as activators or inhibitors of MMP-13 or of an MMP-13 analogue.

More specifically, the present invention provides a method for determining the molecular structure of a molecule or molecular complex whose structure is unknown, comprising the steps of obtaining crystals or a solution of the molecule or molecular complex whose structure is unknown, and then generating x-ray diffraction data from the crystallized molecule or molecular complex, and/or generating NMR data from the solution of the molecule or molecular complex. The x-ray diffraction data from the molecule or molecular complex whose structure is unknown is then compared to the x-ray diffraction data obtained from the MMP-13:Compound A crystal of the present invention.

Alternatively, the NMR data from the molecule or molecular structure whose structure is unknown is then compared with the NMR data obtained from the MMP-13:Compound A solution of the present invention. Then, molecular replacement analysis is used to conform the three dimensional structure determined from the MMP-13:Compound A crystal of solution of the present invention to the x-ray diffraction data from the unknown molecule or molecular complex, or, alternatively, 2D, 3D and 4D isotope filtering, editing and triple resonance NMR techniques are used to conform the three dimensional structure determined from the MMP-13:Compound A solution of the present invention to the NMR data from the solution molecule or molecular complex.

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Molecular replacement analysis uses a molecule having a known structure as a starting point to model the structure of an unknown crystalline sample. This technique is based on the principle that two molecules which have similar structures, orientations and positions will diffract x-rays similarly. A corresponding approach to molecular replacement is applicable to modeling an unknown solution structure using NMR technology. The NMR spectra and resulting analysis of the NMR data for two similar structures will be essentially identical for regions of the proteins that are structurally conserved, where the NMR analysis consists of obtaining the NMR resonance assignments and the structural constraint assignments, which may contain hydrogen bond, distance, dihedral angle, coupling constant, chemical shift and dipolar coupling constant constraints. The observed differences in the NMR spectra of the two structures will highlight the differences between the two structures and identify the corresponding differences in the structural constraints. The structure determination process for the unknown structure is then based on modifying the NMR constraints from the known structure to be consistent with the observed spectral differences between the NMR spectra.

Accordingly, in one non-limiting embodiment of the invention, the resonance assignments for the MMP-13:Compound A complex provide the starting point for resonance assignments of MMP-13 in a new MMP-13:"unsolved agent" complex. Chemical shift perturbances in two dimensional

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¹⁵N/¹H spectra can be observed and compared between the MMP-13:Compound A complex and the new MMP-13:agent complex. In this way, the affected residues may be correlated with the three dimensional structure of MMP-13 as provided by the relevant residues of Figure 4. This effectively identifies the region of the MMP-13:agent complex that has incurred a structural change relative to the MMP-13:Compound A complex. The ¹H, ¹⁵N, ¹³C and ¹³CO NMR resonance assignments corresponding to both the sequential backbone and sidechain amino acid assignments of MMP-13 may then be obtained and the three dimensional structure of the new MMP-13:agent complex may be generated using standard 2D, 3D and 4D triple resonance NMR techniques and NMR assignment methodology, using the MMP-13:Compound A structure, resonance assignments and structural constraints as a reference. Various computer fitting analyses of the new agent with the three dimensional model of MMP-13 may be performed in order to generate an initial three dimensional model of the new agent complexed with MMP-13, and the resulting three dimensional model may be refined using standard experimental constraints and energy minimization techniques in order to position and orient the new agent in association with the three dimensional structure of MMP-13.

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The present invention further provides that the structural coordinates of the present invention may be used with standard homology modeling techniques in order to determine the unknown three-dimensional structure of a molecule or molecular complex. Homology modeling involves constructing a model of an unknown structure using structural coordinates of one or more related protein molecules, molecular complexes or parts thereof (i.e., active sites). Homology modeling may be conducted by fitting common or homologous portions of the protein whose three dimensional structure is to be solved to the three dimensional structure of homologous structural elements in the known molecule, specifically using the relevant (i.e., homologous) structural coordinates provided by Figures 4 and/or 5 herein. Homology may be determined using amino acid sequence identity, homologous secondary structure elements, and/or homologous tertiary folds. Homology modeling can

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include rebuilding part or all of a three dimensional structure with replacement of amino acids (or other components) by those of the related structure to be solved.

Accordingly, a three dimensional structure for the unknown molecule or molecular complex may be generated using the three dimensional structure of the MMP-13:Compound A complex of the present invention, refined using a number of techniques well known in the art, and then used in the same fashion as the structural coordinates of the present invention, for instance, in applications involving molecular replacement analysis, homology modeling, and rational drug design.

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Determination of the three dimensional structure of MMP-13 and its catalytic active site as disclosed herein is critical to the rational identification and/or design of therapeutic agents that may act as inhibitors or activators of MMP-13 enzymatic activity. Alternatively, using conventional drug assay techniques, the only way to identify such an agent is to screen thousands of test compounds, either in culture or by administration to suitable animal models in a laboratory setting, until an agent having the desired inhibitory or activating effect on a target compound is identified. Necessarily, such conventional screening methods are expensive, time consuming, and do not elucidate the method of action of the identified agent on the target compound.

However, advancing X-ray, spectroscopic and computer modeling technologies allow researchers to visualize the three dimensional structure of a targeted compound. Using such a three dimensional structure, researchers identify putative binding sites and then identify or design agents to interact with these binding sites. These agents are then screened for an activating or inhibitory effect upon the target molecule. In this manner, not only are the number of agents to be screened for the desired activity greatly reduced, but the mechanism of action on the target compound is better understood.

Accordingly, the present invention further provides a method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of using a three dimensional structure of MMP-13 as defined by the relative

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structural coordinates of amino acids encoding MMP-13 to design or select a potential inhibitor or activator, and synthesizing or obtaining said potential inhibitor or activator. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of an empty MMP-13 active site in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors. The method of the present invention is preferably used to design or select inhibitors of MMP-13 activity.

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An agent that interacts or associates with an active site of MMP-13 or an MMP-13 analogue may be identified by determining an active site of MMP-13 or of the MMP-13 analogue from a three dimensional model of the MMP-13 or MMP-13 analogue, and performing computer fitting analyses to identify an agent which interacts or associates with said active site. Computer fitting analyses utilize various computer software programs that evaluate the "fit" between the putative active site and the identified agent, by (a) generating a three dimensional model of the putative active site of a molecule or molecular complex using homology modeling or the atomic structural coordinates of the active site, and (b) determining the degree of association between the putative active site and the identified agent. The degree of association may be determined computationally by any number of commercially available software programs, or may be determined experimentally using standard binding assays.

Three dimensional models of the putative active site may be generated using any one of a number of methods known in the art, and include, but are not limited to, homology modeling as well as computer analysis of raw structural coordinate data generated using crystallographic or spectroscopy techniques. Computer programs used to generate such three dimensional models and/or perform the necessary fitting analyses include, but are not limited to: GRID (Oxford University, Oxford, UK), MCSS (Molecular Simulations, San Diego, CA), AUTODOCK (Scripps Research Institute, La Jolla,

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CA), DOCK (University of California, San Francisco, CA), Flo99 (Thistlesoft, Morris Township, NJ), Ludi (Molecular Simulations, San Diego, CA), QUANTA (Molecular Simulations, San Diego, CA), Insight (Molecular Simulations, San Diego, CA), SYBYL (TRIPOS, Inc., St. Louis. MO) and LEAPFROG (TRIPOS, Inc., St. Louis, MO).

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In a preferred method of the present invention, the identified active site of MMP-13 or the MMP-13 analogue comprises a catalytic zinc, a beta strand, a Ca²⁺ binding loop, an alpha helix and a random coil region. More preferably, the identified active site comprises a catalytic zinc, a beta strand comprising residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1 (or conservative substitutions thereof), a Ca²⁺ binding loop comprising residues F75, D76, G77, P78, and S79 according to Figure 1 (or conservative substitutions thereof), an alpha helix comprising residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1 (or conservative substitutions thereof), and a random coil region comprising residues P139, I140, and Y141 according to Figure 1 (or conservative substitutions thereof).

More specifically, the identified active site of the present method comprises the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). In an alternate preferred embodiment, the identified active site further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case,

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± a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). In yet a third preferred embodiment, the identified active site of the present method further comprises the relative structural coordinates of amino acid residues F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). Embodiments comprising conservative substitutions of the noted amino acids result in the same structural coordinates of the corresponding residues in Figures 4 or 5 within the stated root mean square deviation.

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The effect of such an agent identified by computer fitting analyses on MMP-13 (or MMP-13 analogue) activity may be further evaluated computationally, or experimentally by contacting the identified agent with MMP-13 (or an MMP-13 analogue) and measuring the effect of the agent on the enzyme's activity. Depending upon the action of the agent on the active site of MMP-13, the agent may act either as an inhibitor or activator of MMP-13 activity. Standard enzymatic assays may be performed and the results analyzed to determine whether the agent is an inhibitor of MMP-13 activity (i.e., the agent may reduce or prevent binding affinity between MMP-13 and the relevant substrate, and thereby reduce the level or rate of MMP-13 activity compared to baseline), or an activator of MMP-13 activity (i.e., the agent may increase binding affinity between MMP-13 and the relevant substrate, and thereby increase the level or rate of MMP-13 activity compared to baseline). Further tests may be performed to evaluate the selectivity of the identified agent to MMP-13 with regard to the other metalloproteinases.

Agents designed or selected to interact with MMP-13 must be capable of both physically and structurally associating with MMP-13 via various covalent and/or non-covalent molecular interactions, and of assuming a three

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dimensional configuration and orientation that complements the relevant active site of the MMP-13 molecule.

Accordingly, using these criteria, the structural coordinates of the MMP-13:Compound A complex as disclosed herein, and/or structural coordinates derived therefrom using molecular replacement analysis or homology modeling, agents may be designed to increase either or both of the potency and selectivity of known inhibitors or activators, either by modifying the structure of known inhibitors or activators or by designing new agents *de novo* via computational inspection of the three dimensional configuration and electrostatic potential of an MMP-13 active site.

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Accordingly, in one embodiment of the invention, the structural coordinates of Figures 4 or 5 of the present invention, or structural coordinates derived therefrom using molecular replacement or homology modeling techniques as discussed above, are used to screen a database for agents that may act as potential inhibitors or activators of MMP-13 activity (or the activity of MMP-13 analogues). Specifically, the obtained structural coordinates of the present invention are read into a software package and the three dimensional structure is analyzed graphically. A number of computational software packages may be used for the analysis of structural coordinates, including, but not limited to, Sybyl (Tripos Associates), QUANTA and XPLOR (Brunger, A.T., (1993) XPLOR Version 3.1 Manual, Yale University, New Haven, CT). Additional software programs check for the correctness of the coordinates with regard to features such as bond and atom types. If necessary, the three dimensional structure is modified and then energy minimized using the appropriate software until all of the structural parameters are at their equilibrium/optimal values. The energy minimized structure is superimposed against the original structure to make sure there are no significant deviations between the original and the energy minimized coordinates.

The energy minimized coordinates of MMP-13 complexed with a "solved" inhibitor or activator are then analyzed and the interactions between the solved ligand and MMP-13 are identified. The final MMP-13 structure is

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modified by graphically removing the solved inhibitor or activator so that only MMP-13 and a few residues of the solved agent are left for analysis of the binding site cavity. QSAR and SAR analysis and/or conformational analysis may be carried out to determine how other inhibitors or activators compare to the solved inhibitor or activator. The solved agent may be docked into the uncomplexed structure's binding site to be used as a template for data base searching, using software to create excluded volume and distance restrained queries for the searches. Structures qualifying as hits are then screened for activity using standard assays and other methods known in the art.

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Further, once the specific interaction is determined between the solved inhibitor or activator, docking studies with different inhibitors or activators allow for the generation of initial models of new inhibitors or activators in complex with MMP-13. The integrity of these new models may be evaluated a number of ways, including constrained conformational analysis using molecular dynamics methods (*i.e.*, where both MMP-13 and the complexed activator or inhibitor are allowed to sample different three dimensional conformational states until the most favorable state is reached or found to exist between the protein and the complexed agent). The final structure as proposed by the molecular dynamics analysis is analyzed visually to make sure that the model is in accord with known experimental SAR based on measured binding affinities. Once models are obtained of the original solved agent bound to MMP-13 and computer models of other molecules bound to MMP-13, strategies are determined for designing modifications into the activators or inhibitors to improve their activity and/or enhance their selectivity.

Once an MMP-13 binding agent has been optimally selected or designed, as described above, substitutions may then be made in some of its atoms or side groups in order to improve or modify its selectivity and binding properties. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the

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original group. Such substituted chemical compounds may then be analyzed for efficiency of fit to MMP-13 by the same computer methods described in detail above.

Alternatively, the present invention provides a method for identifying a potential inhibitor or activator that is selective for one or more members of the matrix metalloproteinase family except MMP-13, comprising the steps of (i) using the three dimensional structures of MMP-13 and the desired target matrix metalloproteinase(s) as defined by the relative structural coordinates of amino acids encoding MMP-13 and the target matrix metalloproteinase(s) in order to design or select such a potential inhibitor or 10 activator, and (ii) synthesizing or obtaining said potential inhibitor or activator. In this case, the potential inhibitor or activator is designed to incorporate chemical or steric features favorable for association with an active site of the desired matrix metalloproteinase(s) and unfavorable for association with an MMP-13 active site, preferably where said active site comprises the MMP-13 S1' pocket. The inhibitor or activator may be selected by screening an appropriate database, may designed de novo by analyzing the steric configurations and charge potentials of empty MMP-13/matrix metalloproteinase active sites in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other 20 collagenases in order to create "hybrid" activators or inhibitors.

Various molecular analysis and rational drug design techniques are further disclosed in U.S. Patent Nos. 5,834,228, 5,939,528 and 5,865,116, as well as in PCT Application No. PCT/US98/16879, published as WO 99/09148, the contents of which are hereby incorporated by reference.

The present invention may be better understood by reference to the following non-limiting Examples. The following Examples are presented in order to more fully illustrate the preferred embodiments of the invention, and should in no way be construed as limiting the scope of the present invention.

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Example 1

¹H, ¹⁵N and ¹³CO Assignments and Secondary Structure Determination of MMP-13 Complexed with Compound A

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Methods and Results: The uniform ¹⁵N and ¹³C- labeled 165 amino-acid catalytic fragment of human collagenase-3 (MMP-13) was expressed in *E. coli* strain BL21(DE3) containing the plasmid pProMMP-13 according to a published method (Freije *et al.*, J. <u>Biol. Chem.</u> 1994). MMP-13 was purified as previously described (Moy *et al.*, J. <u>Biomol.</u> 1997) with minor modifications. N-terminal amino acid sequencing was performed to confirm the protein's identity while the uniform ¹⁵N and ¹³C labeling of MMP-13 was confirmed by MALDI-TOF mass spectrometry (PerSeptive Biosystems). The sulfonamide derivative of the hydroxamic acid compound, N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide, was prepared from 2-amino-3-methyl-benzoic acid methyl ester and p-methoxybenzenesulfonyl chloride followed by alkylation with 3-picolyl chloride, hydrolysis (LiOH/THF) to afford the carboxylic acid and conversion to the hydroxamic acid (oxalyl chloride/DMF/NH2OH). Formation of the HCl salt yielded Compound A as shown in Figure 3.

The NMR samples contained 1 mM of MMP-13 determined spectrophotometrically in a equimolar complex with Compound A in a buffer containing 10 mM deuterated Tris-Base, 100 mM NaCl, 5 mM CaCl₂, 0.1 mM $\rm ZnCl_2$, 2 mM NaN₃, 10 mM deuterated DTT, in either 90% $\rm H_2O/$ 10% $\rm D_2O$ or 100% $\rm D_2O$ at pH 6.5. All NMR spectra were recorded at 35°C on a Bruker AMX-2 600 spectrometer equipped with a triple-resonance gradient probe.

Spectra were processed using the NMRPipe software package (Delaglio *et al.*, <u>J. Biomol. NMR</u> 1995) and analyzed with PIPP (Garrett *et al.*, <u>J. Magn. Reson.</u> 1991), NMRPipe and PEAK-SORT, an in-house software package. The assignments of the ¹H, ¹⁵N, ¹³CO, and ¹³C resonances were based on the following experiments: CBCA(CO)NH, CBCANH, C(CO)NH, HC(CO)NH,

HBHA(CO)NH, HNCO, HCACO, HNHA, HNCA, HCCH-COSY and HCCH-TOCSY (for reviews, see Bax *et al.*, Methods Enzymol. 1994; and Clore & Gronenborn, Methods Enzymol. 1994). The accuracy of the MMP-13 NMR assignments was further confirmed by sequential NOEs in the ¹⁵N-edited NOESY-HSQC spectra.

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Prior to analysis of the MMP-13 NMR structure, the structure determination of the inhibitor-free catalytic fragment of MMP-1 has been reported (Moy et al., Biochemistry 1998; Moy et al., J. Biomol. NMR 1997)(30 simulated annealing structures deposited with Protein Data Bank, Accession No. 1AYK; restrained minimized mean structure deposited with Protein Data Bank, Accession No. 2AYK). Because the MMPs are highly autocatalytic, the NMR analysis of the inhibitor-free MMP-1 was accomplished by establishing buffer conditions where the enzyme was still active but the rate of self-cleavage of the enzyme had been diminished. This was achieved by the addition of DTT which significantly diminished self-aggregation of the enzyme and by lowering the pH of the sample to 6.5, just above the pH where the enzyme was known to be inactivated because of the loss of the catalytic zinc. Under these conditions, an MMP-1 NMR sample was typically stable for 1-2 months. Unfortunately this was not the case for MMP-13, the protein rapidly degraded within a few hours which required the use of an inhibitor to assign the MMP-13 NMR resonances.

The secondary structure of the MMP-13:Compound A complex is based on characteristic NOE data involving the NH, H α and H β protons from 15 N-edited NOESY-HSQC and 13 C-edited NOESY-HMQC spectra, 3 JHN α coupling constants from HNHA, slowly exchanging NH protons and 13 C α and 13 C β secondary chemical shifts (for reviews, see Wishart & Sykes, Methods Enzymol. 1994; and Wuthrich, NMR of Proteins and Nucleic Acids, John Wiley & Sons, New York 1986). It was determined that the MMP-13 NMR structure in the complex is composed of three α -helices corresponding to residues 28-44 (a $_{\alpha}$), 112-123 (a $_{\beta}$) and 153-163 (a $_{c}$) and a mixed parallel and anti-parallel β -sheet consisting of 5 strands corresponding to residues 83-86 (β_{1}), 95-100 (β_{2}), 59-66 (β_{3}), 14-20 (β_{4}) and 49-53 (β_{5}). This is essentially identical to the secondary structure observed for other MMP structures.

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There were three distinct regions in the MMP-13:Compound A spectra where the resonance assignments are incomplete. These correspond to residues G70-Y73, P87-N91 and T144-H148. Residues T144-H148 correspond to part of the dynamic loop region previously seen in the MMP-1 structure (Moy et al., J. Biomol. NMR 1997). This suggests a similar dynamic profile for this region in the MMP-13 structure even in the presence of a high-affinity inhibitor (IC₅₀ = 33 nM). Residues P87 to N91 contain a cluster of prolines which disrupt the sequential assignment process because of the missing NH. Residues G70 to Y73 correspond to a loop region in the vicinity of the structural zinc which was readily assigned in the MMP-1 structure. The backbone and side-chain 1 H, 15 N, 13 C, and 13 CO assignments are essentially complete for the remainder of the protein.

Example 2

High Resolution Solution Structure of the Catalytic Fragment of MMP-13
Complexed with Compound A

Materials and Methods:

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Preparation of Compound A: The sulfonamide derivative of the hydroxamic acid compound, Compound A, was prepared according to the procedure noted in Example 1 to yield the compound of Figure 3.

Expression of recombinant ¹⁵N and ¹³C/ ¹⁵N-labeled MMP-13: A 169-residue C-terminally truncated human collagenase-3 (MMP-13) was expressed in *E. coli*.

The coding sequence of a C-terminally truncated procollagenase was amplified by PCR from the plasmid pNot3a, that contains the entire coding sequence of MMP-13 (Frieje, *et al.*, J. <u>Biol. Chem.</u> 1994). The PCR primers contained the appropriate restriction sites for ease of cloning. The construct codes for a truncated proMMP-13 with an N-terminal methionine added and a C-terminal proline at residue 169 of the native proMMP-13 sequence. The PCR amplified DNA fragment was the cloned into pET-21a (+) at the Nde I/Sal I sites,

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resulting in a recombinant plasmid designated as pProMMP-13. *E. coli* bacteria, BL21(DE3), containing the plasmid pProMMP-13, were grown in LB broth supplemented with 100 μ g/ml ampicilin. An overnight culture was diluted 1:20 and grown at 37°C to an A₆₀₀ of 0.6-0.8 with vigorous shaking. Isopropyl β -D-galactoside (IPTG) was added to a final concentration of 1 mM and cultures were shaken for 3 h at 37°C. The cells were harvested by centrifugation (7000 Xg for 15 min) at 4°C, washed with PBS, and frozen at -70°C until further use.

Uniform 15 N and 13 C- labeled ProMMP-13 was obtained by growing BL21(DE3) E. coli in defined media containing 2.0 g/l [13 C6, 98%+]D-glucose and 1.0 g/l [15 N, 98%+] ammonium chloride as the sole carbon and nitrogen sources, respectively. In addition, the defined media contained M9 salts (Sambrook, *et al.*, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, New York, NY 1989), trace elements, vitamins and 100 μ g/ml ampicilin. Conditions for induction and growth were the same as above.

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Purification of recombinant ¹⁵N and ¹³C MMP-13: MMP-13 was purified according to Moy et al., J. Biomol. NMR 1997, with modifications as follows. Frozen cell pellets were thawed on ice. Cells were resuspended by homogenization in lysis buffer (0.1 M Tricine, pH 8.0, 10 mM EDTA, 2mM DTT, 0.5 mM PMSF). Cells 20 were lysed by French Press (2X) followed by treatment with lysozyme (1 mg/ml; final) at room temperature for 30 min. The lysate was centrifuged at 45,000 x g for 30 minutes. The pellet was washed twice with 50 mM Tricine pH 7.5, 0.2 M NaCl₂, 0.5% Triton X-100, resuspended in fresh urea buffer (20 mM Tricine, pH 7.5, 8 M urea, 0.2% NaN₃, 2 mM DTT) and incubated at room temperature for I hour. The urea solubilized protein was centrifuged at 45,000 x g for 30 min and the resultant supernatant was filtered and applied to a Hitrap-Q Sepharose (Pharmacia Biotech) anion exchange column equilibrated in 6 M urea buffer. The column was washed with urea buffer and eluted with a 0-0.25 M NaCl linear gradient. Fractions containing proMMP-13 were detected by 30 SDS-PAGE, pooled and quickly diluted into 5-fold excess of renaturing buffer

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(50 mM Tricine, pH 7.5, 0.4 M NaCl, 10 mM CaCl₂, 0.1 mM ZnOAc₂, 0.02% NaN₃). After 2 days of dialysis against 25 volumes of renaturing buffer (with three changes), refolded proMMP-13 was concentrated to about 4-10 mg/ml in a Millipore Biomax 5 concentrator. ProMMP-13 was activated to MMP-13CAT (catalytic domain) by an overnight incubation at 37 °C in the presence of l mM p-aminophenylmercuric acetate (APMA).

The activated protein is then applied onto a Superdex-75 16/60 gel filtration column equilibrated in 2.5 mM Tris-HCl, pH 7.5, 5 mM CaCl₂, 0.4 M NaCl, 2 mM DTT, 0.02% NaN₃ and 0.05 mM ZnOAc₂. The protein is eluted and fractions containing MMP-13CAT were identified by SDS-PAGE. Peak fractions were pooled and the protein was concentrated in a Millipore Biomax concentrator to about 5 mg/ml and stored at -70 °C. N-terminal amino acid sequencing was performed to confirm the protein's identity. The uniform ¹⁵N and ¹³C labeling of MMP-13-CAT was confirmed by MALDI-TOF mass spectrometry (PerSeptive Biosystems).

NMR Sample Preparation: The MMP-13:Compound A NMR sample contained $1\,\mathrm{mM}^{15}\mathrm{N}$ -or $^{15}\mathrm{N}/^{13}\mathrm{C}$ -labeled MMP-13 with Compound A in a 1:1 ratio. The sample was prepared by repeated buffer exchange using 20-30ml solution containing 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl₂, 0.1mM ZnCl₂, 2mM NaN₃, 10mM deuterated DTT, and 0.2mM Compound A in either 90% H₂O/10 % D₂O or 100% D₂O. Buffer exchange was carried out on a Millipore Ultrafree-15 Centrifugal Filter Unit. Excess Compound A was removed by additional buffer exchanges where Compound A was removed from the buffer.

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NMR Data Collection: All spectra were recorded at 35°C on a Bruker AMX-2 600 spectrometer using a gradient enhanced triple-resonance ¹H/¹³C/¹⁵N probe. For spectra recorded in H₂O, water suppression was achieved with the WATERGATE sequence and water-flip back pulses (Piotto, et al., J. <u>Biomol. NMR</u> 1992; Grzesiek and Bax, J. <u>Am. Chem. Soc.</u> 1993). Quadrature detection in the

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indirectly detected dimensions were recorded with States-TPPI hypercomplex phase increment (Marion, et al., J. Magn. Reson. 1989). Spectra were collected with appropriate refocusing delays to allow for 0,0 or -90,180 phase correction.

The resonance assignments and bound conformation of Compound A in the MMP-1: Compound A complex were based on the 2D ¹²C/¹²C-filtered NOESY (Petros, et al., <u>FEBS Lett.</u> 1992; Gemmecker, et al., <u>J. Magn. Reson.</u> 1992), 2D ¹²C/¹²C-filtered TOCSY (Petros, et al., <u>FEBS Lett.</u> 1992; Gemmecker, et al., <u>J. Magn. Reson.</u> 1992) and ¹²C/¹²C-filtered COSY experiments (Ikura and Bax, <u>J. Magn. Reson.</u> 1992).

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The MMP-13:Compound A structure is based on the following series of spectra: HNHA (Vuister and Bax, J. Am. Chem. Soc. 1993), HNHB (Archer, et al., J. Magn. Reson. 1992), 3D long-range ¹³C-¹³C correlation (Bax and Popchapsky, J. Magn. Reson. 1992), coupled CT-HCACO (Powers, et al., J. Magn. Reson. 1991; Vuister, et al., J. Am. Chem. Soc. 1992), HACAHB-COSY (Grzesiek, et al., J. Amer. Chem. Soc. 1995), 3D ¹⁵N- (Mario, et al., Biochemistry 1989; Zuiderweg and Fesik, Biochemistry 1989) and ¹³C-edited NOESY (Zuiderweg, et al., J. Magn. Reson. 1990; Ikura, et al., J. Magn. Reson. 1990), and 3D ¹³C-edited/¹²C-filtered NOESY (Lee, et al., FEBS Lett. 1994). experiments. The ¹⁵N-edited NOESY, ¹³C-edited NOESY and 3D ¹³C-edited/¹²C-filtered NOESY experiments were collected with 100 msec, 120 msec and 110 msec mixing times, respectively. The acquisition parameters for each of the experiments used in determining the solution structure of MMP-13 complexed with Compound A were as reported previously (Moy, et al., Biochemistry, 1998).

Spectra were processed using the NMRPipe software package (Delaglio, et al., J. Biomol. NMR, 1995) and analyzed with PIPP (Garrett, et al., J. Magn. Reson., 1991) on a Sun Sparc Workstation. When appropriate, data processing included a solvent filter, zero-padding data to a power of two, linear predicting back one data point of indirectly acquired data to obtain zero phase corrections, linear prediction of additional points for the indirectly acquired dimensions to increase resolution. Linear prediction by the means of the mirror

image technique was used only for constant-time experiments (Zhu and Bax, J. Magn. Reson., 1992). In all cases data was processed with a skewed sine-bell apodization function and one zero-filling was used in all dimensions.

- 5 Interproton Distance Restraints: The NOEs assigned from 3D ¹³C-edited/¹²C-filtered NOESY and 3D ¹⁵N-edited NOESY experiments were classified into strong, medium, and weak corresponding to interproton distance restraints of 1.8-2.7 Å (1.8-2.9 Å for NOEs involving NH protons), 1.8-3.3 Å (1.8-3.5 Å for NOEs involving NH protons), and 1.8-5.0 Å, respectively (Williamson, et al., J. Mol. Biol., 1985; Clore, et al., EMBO J., 1986). Upper distance limits for
- Mol. Biol., 1985; Clore, et al., EMBO J., 1986). Upper distance limits for distances involving methyl protons and non-stereospecifically assigned methylene protons were corrected appropriately for center averaging (Wuthrich, et al., J. Mol. Biol., 1983).
- Torsion Angle Restraints and Stereospecific Assignments. The β-methylene stereospecific assignments and χ₁ torsion angle restraints were obtained primarily from a qualitative estimate of the magnitude of ³J_{αβ} coupling constants from the HACAHB-COSY experiment (Grzesiek, et al., J. Am. Chem. Soc., 1992) and ³J_{Nβ} coupling constants from the HNHB experiment (Archer, et al., J. Magn.
- Reson., 1991). Further support for the assignments was obtained from approximate distance restraints for intraresidue NOEs involving NH, C α H, and C β H protons (Powers, et al., Biochemistry, 1993).

The ϕ and ψ torsion angle restraints were obtained from ${}^3J_{NH\alpha}$ coupling constants measured from the relative intensity of H α crosspeaks to the NH diagonal in the HNHA experiment (Vuister and Bax, J. Am. Chem. Soc. 1993), from a qualitative estimate of the magnitude of ${}^3J_{\alpha\beta}$ coupling constants from the HACAHB-COSY experiment (Grzesiek, et al., J. Am. Chem. Soc., 1992) and from approximate distance restraints for intraresidue and sequential NOEs involving NH, C α H, and C β H protons by means of the conformational grid search program STEREOSEARCH (Nilges, et al., Biopolymers 1990), as described previously (Kraulis, et al., Biochemistry 1989). ${}^1J_{c\alpha H\alpha}$ coupling

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constants obtained from a coupled 3D CT-HCACO spectrum were used to ascertain the presence of non-glycine residues with positive f backbone torsion angles (Vuister, et al., J. Am. Chem. Soc. 1992). The presence of a $^1J_{c\alpha H\alpha}$ coupling constant greater then 130 Hz allowed for a minimum φ restraint of -2° to -178°.

The Ile and Leu χ2 torsion angle restraints and the stereospecific assignments for leucine methyl groups were determined from ${}^3J_{\text{CαCδ}}$ coupling constants obtained from the relative intensity of Cα and Cδ cross peaks in a 3D long-range ${}^{13}\text{C}$ - ${}^{13}\text{C}$ NMR correlation spectrum (Bax, et al., J. Am. Chem. Soc. 1992), in conjunction with the relative intensities of intraresidue NOEs (Powers, et al., Biochemistry 1993). Stereospecific assignments for valine methyl groups were determined based on the relative intensity of intraresidue NH-CγH and CαH-CγH NOEs as described by Zuiderweg et al. (1985) (Zuiderweg, et al., Biopolymers 1985). The minimum ranges employed for the φ , ψ , and χ torsion angle restraints were \pm 30°, \pm 50°, and \pm 20°, respectively (Kraulis, et al., Biochemistry 1989).

Structure Calculations: The structures were calculated using the hybrid distance geometry-dynamical simulated annealing method of Nilges et al. (1988)

20 (Protein Eng.) with minor modifications (Clore, et al., Biochemistry 1990) using the program XPLOR (Brunger, X-Plor Version 3.1 Manual, Yale University, New Haven, CT), adapted to incorporate pseudopotentials for ³J_{NHα} coupling constants (Garrett, et al., J. Magn. Reson. Ser. B 1994), secondary ¹³Cα/¹³Cβ chemical shift restraints (Kuszewski, et al., J. Magn. Reson. Ser B 1995) and a conformational database potential (Kuszewski, et al., Protein Sci. 1996; Kuszewski, et al., J. Magn. Reson. 1997). The target function that is minimized during restrained minimization and simulated annealing comprises only quadratic harmonic terms for covalent geometry, ³J_{NHα} coupling constants and secondary ¹³Cα/¹³Cβ chemical shift restraints, square-well quadratic potentials for the experimental distance and torsion angle restraints, and a quartic van der

Waals term for non-bonded contacts. All peptide bonds were constrained to be

planar and trans. There were no hydrogen-bonding, electrostatic, or 6-12 Lennard-Jones empirical potential energy terms in the target function.

To prevent the Zn and Ca ions from being expelled during the high-temperature simulated annealing stages of the refinement protocol, a minimal number of distance restraints between the His sidechain and Zn and between backbone atoms and Cα were included in the XPLOR distance restraint file based on the observed coordination in the X-ray structures (Lovejoy, *et al.*, Science 1994; Lovejoy, *et al.*, Biochemistry 1994; Spurlino, *et al.*, Proteins: Struct., Funct., Genet. 1994; Borkakoti, *et al.*, Nat. Struct. Biol. 1994).

The starting MMP-13:Compound A complex structure for the simulated-annealing protocol was obtained by manually docking Compound A into a homology model for MMP-13. The initial orientation of Compound A in the MMP-13 active site was based on the previously reported MMP-1:CGS-27023A structure (Moy, et al., Biochemistry 1999).

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Homology modeling methods were utilized to generate a three dimensional model of MMP-13. The linear amino acid sequence corresponding to the catalytic domain of MMP-13 was aligned (SYBYL) with the catalytic domains of MMP-1, MMP-7 and MMP-8 based on the availability of their x-ray crystallographic structures (Bode, et al., EMBO J 1994; Spurlino., Proteins: Struct., Funct., Genet. 1994; Betz, et al., Eur. J. Biochem. 1997; Lovejoy, et al., Nat. Struct. Biol. 1999; Borkakoti, et al., Nat. Struct. Biol. 1994; Browner, et al., Biochemistry 1995). The alignments of MMP-13 with MMP-1 and MMP-8 demonstrated the highest homology where the computed identities are 58.9% and 61.4%, respectively (Figure 2).

The X-ray structure of MMP-8 was selected to be used as the template for homology modeling the structure of MMP-13. This decision was based mainly on the sequence alignment shown in Figure 2B where no insertions (labeled "###") are found in the critical specificity loop (Labeled Underlined and Boldface). In Figure 2A, the region labeled "##" in the specificity loop shows that there is an "insertion" of 2 additional amino acid residues compared to the sequence length of MMP-1. Based on our analysis of

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the alignments, MMP-8 would allow for a more accurate modeling of the inhibitor binding pockets since no predictions have to be made within this loop region.

Tomposer (SYBYL) was used to construct the initial homology model of MMP-13. The only insertion was a serine (labeled '**' in Figure 2B) at position 32 of MMP-13. The insertion of S32 occurs within a coiled region which is at the entrance of a long alpha helix and about 17 angstroms from the S' specificity loop. The model of MMP-13 was then energy minimized utilizing a set of nested refinement procedures (Chen, et al., J. Biomol. Struct. Dyn. 1995), but where the protein backbone heavy atoms were constrained as close as possible to their original positions.

The MMP-13:Compound A model was then subjected to a 1000 steps of CHARMM minimization with the 5 intramolecular NOE restraints and the 47 distance restraints observed between MMP-13 and Compound A where the coordinates for MMP-13 were kept fixed. This approach approximated the positioning of Compound A in the active site of MMP-13 without distorting the MMP-13 structure. The final structure was exported as a PDB file and used as the starting point for XPLOR simulated annealing protocol where all the residues in the structure were free to move. Since the initial stage of the simulated annealing protocol corresponds to high-temperature dynamics (1500 K) with a relatively weak XPLOR NOE force constant (Ries and Petrides, Biol. Chem. Hoppe-Seyler 1995), the initial MMP-13:Compound A structure does not bias the structure determination process since the structure is effectively free to explore the available conformational space. Additionally, each iteration of the simulated annealing process begins with a random trajectory for the molecular dynamics. The fact that these trajectories differ by upwards of 10 Å assures a distinct exploration of conformational space for the ensemble of MMP-13:Compound A structures determined from the simulated annealing protocol.

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Results and Discussion

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Compound A Resonance Assignments and Bound Conformation: The primary structure of Compound A along with the proton naming convention is shown in Figure 3. The NMR assignments for Compound A in the MMP-13 complex followed established protocols using 2D ¹²C-filtering experiments (Petros, et al., FEBS Lett. 1992; Gemmecker, et al., J. Magn. Reson. 1992; Ikura and Bax, J. Am. Chem. Soc. 1992) since the NMR sample was composed of ¹³C/¹⁵N labeled MMP-13 and unlabeled Compound A. Thus, traditional 2D-NOESY, COSY and TOCSY spectra of Compound A in the presence of MMP-13 yielded straightforward assignments for Compound A along with assignments for free 10 Compound A (data not shown). The only notable difference in the assignments for free and bound Compound A is the observation of two distinct resonances for 2HB1/2 in the complex (4.91 ppm; 4.67 ppm). The missing resonance in the free Compound A may simply be obscured by water. Also, an observation that the protons on the p-methoxyphenyl ring are degenerate suggests rapid ring flips when complexed to MMP-13. This was also seen with CGS-27023A complexed with both MMP-1 and stromelysin (Gonnella, et al., Bioorg. Med. Chem. 1997; Moy, et al., Biochemistry 1998; Moy, et al., Biochemistry 1999).

absence of MMP-13 as evident by the lack of structural NOEs. Only a minimal number of intramolecular NOEs were observed for Compound A in the MMP-13 complex which were relevant to the bound conformation of Compound A (data not shown). The minimal number of structural NOEs is a result of the Compound A conformation, structure and chemical shift degeneracy. A number of the observed NOEs correspond to a sequential interaction which have no effect on the overall conformation of the inhibitor and were not used in the refinement of Compound A or the complex. The structural intramolecular NOEs observed are primarily between 1HH* and the pyridine ring and between 2HB1/2 and both the p-methoxyphenyl and aryl ring. These NOEs are consistent with the "splayed" conformation previously observed for CGS-27023A bound to both MMP-1 and stromelysin, but the bound conformation of

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Compound A is predominately determined from the intermolecular NOEs between Compound A and MMP-13 (Table 1).

Structure Determination: The NMR structure determination methodology is an
iterative procedure where the current state of the structure is used to analyze the ambiguous NOE data. In essence, the structure is used as a distance filter to sort through the ambiguous NOE list where the first structure is determined from unambiguous data. For the refinement of MMP-13, the initial structure was a homology model based on the MMP-8 X-ray structure. This was justified
by the overall similarity in previously reported MMP structures and from the secondary structure assignments by NMR for MMP-13. The regular secondary structure elements of MMP-13 were identified from a qualitative analysis of sequential and inter-strand NOEs, NH exchange rates, ³JHNα coupling constants (Clore, et al., Crit. Rev. Biochem. Mol. Biol. 1989) and the ¹³Cα and ¹³Cβ
secondary chemical shifts (Spera and Bax, J. Am. Chem. Soc. 1991). The deduced secondary structure is essentially identical to the inhibitor-free MMP-1 NMR structures previously reported.

The final 30 simulated annealing structures calculated for residues 7-164 were based on 3279 experimental NMR restraints, consisting of 2561 approximate interproton distance restraints, 51 distance restraints between MMP-13 and Compound A, 88 distance restraints for 44 backbone hydrogen bonds, 391 torsion angle restraints, $103~^3J_{NH\alpha}$ restraints 123 C α restraints and 108 C β restraints. Stereospecific assignments were obtained for 81 of the 100 residues with β -methylene protons, for the methyl groups of 5 of the 6 Val residues, and for the methyl groups of 12 of the 13 Leu residues. In addition, 12 out of the 12 Phe residues and 7 out of the 8 Tyr residues were well defined making it possible to assign NOE restraints to only one of the pair of C δ H and C ϵ H protons and to assign a χ 2 torsion angle restraint. Similarly, χ 2 torsion angle restraints were assigned for the three Trp residues. The atomic rms distribution of the 30 simulated annealing structures about the mean coordinate positions for residues 7-164 is 0.43 \pm 0.06 Å for the backbone atoms, 0.81 \pm

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0.09 Å for all atoms, and 0.47 \pm 0.04 Å for all atoms excluding disordered surface side chains. The mean standard deviation for the φ and ψ backbone torsion angles of residues 7-164 are 6.2 \pm 11.3° and 7.1 \pm 11.8°, respectively. The high quality of the MMP-13 NMR structure is also evident by the results of PROCHECK analysis and by a calculated, large negative value for the Lennard-Jones-van der Waals energy (-695 \pm 11 kcal mol⁻¹). For the PROCHECK statistics, an overall G-factor of 0.16 \pm 0.16, a hydrogen bond energy of 0.82 \pm 0.05 and only 7.8 \pm 1.0 bad contacts per 100 residues are consistent with a good quality structure comparable to ~1Å X-ray structure.

The high quality of the MMP-13 NMR structure is also evident by the very small deviations from idealized covalent geometry, by the absence of interproton distance and torsion angle violations greater than 0.1 Å and 1°, respectively and by the fact that most of the backbone torsion angles for non-glycine residues lie within expected regions of the Ramachandran plot (not shown). 91.5% of the residues lie within the most favored region of the Ramachandran φ , ψ plot and 7.8% in the additionally allowed regions. 1 JC α H α coupling constants from the coupled CT-HCACO experiment indicated that all non-glycine residues have negative φ torsion angles.

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The quality of the NMR data to properly define the complex is also supported by the well-defined coordinates for Compound A and the active site residues, where the atomic rms distribution is 0.47 ± 0.08 Å and 0.18 ± 0.03 Å for the heavy atoms of Compound A and MMP-13 backbone atoms, respectively.

Description of the MMP-13:Compound A Structure: The overall fold of MMP-13 is
essentially identical to previously reported MMP structures (Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Science 1994; Lovejoy, et al., Ann. N. Y. Acad. Sci. 1994; Lovejoy, et al., Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl. Acad.
Sci. U. S. A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. USA 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr.

1996; Gooley, et al., <u>J. Biomol. NMR</u> 1996; Betz, et al., <u>Eur. J. Biochem.</u> 1997; Gonnella, et al., Bioorg. Med. Chem. 1997; Moy, et al., Biochemistry 1998 and Moy, et al., Biochemistry 1999). The MMP-13 NMR structure is composed of three α -helices corresponding to residues 28-44 (α_A), 112-123 (α_B) and 153-163 (α_c) and a mixed parallel and anti-parallel b-sheet consisting of 5 strands corresponding to residues 83-86 (β_1), 95-100 (β_2), 59-66 (β_3), 14-20 (β_4) and 49-53 (β_5). The active site of MMP-13 is bordered by β -strand IV, the Ca⁺² binding loop, helix B and a random coil region from residues P139-Y141. The catalytic zinc is chelated by H119, H123, and H129 while the structural zinc is chelated by H69, H84 and H97. The calcium ion is chelated in a loop region 10 consisting of residues D75 to G79. An interesting feature of the MMP active-site structure is an apparent kink in the backbone that occurs between the Ca⁺² binding loop and β -strand IV. In the case of MMP-13, this results in the NHs of both L82 and A83 facing toward the active site of the enzyme. An important feature of substrate and inhibitor binding to the MMPs are hydrogen bonding interactions with β -strand IV which is facilitated by this unusual kink conformation (Lovejoy, et al., Science 1994; Lovejoy, et al., Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; and Borkakoti, et al., Nat. Struct. Biol. 1994).

The interaction of Compound A in the active site of MMP-13 was determined by 5 intramolecular NOEs for Compound A and by a total of 47 intermolecular distance restraints between MMP-13 and Compound A. The key MMP-13 residues involved in the interaction with the inhibitor correspond to three distinct MMP-13 regions: residues L81, L82 and A83 from β -strand IV; residues L115, V116, and H119 from α -helix II; and L136, I140 and Y141 from the active site loop which comprise the S1' and S2' pockets of MMP-13. A unique feature of the MMP-13 structure is the large S1' pocket which nearly reaches the surface of the protein.

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Compound A binds to the right-side of the catalytic Zn where the p-methoxyphenyl of Compound A sits in the S1' pocket of the MMP-13 active site. This positioning is evident from the observed NOEs from 3HH*, 3HE1/2

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and 3HD1/2 to L115, V116, H119, L136, and Y141. The aryl group primarily interacts with the side-chain of L81 as evident by the strong NOEs between 1HH*, 1HE2 and 1HZ and the L81 spin-system. Finally, the pyridine ring is essentially solvent exposed but interacts with the side-chain of I140. These interactions position Compound A such that the hydroxamic acid moiety of Compound A chelates to the "right" of the catalytic zinc and the sulfonyl oxygens are in hydrogen-bonding distance to the backbone NH of L82.

It is interesting to note that the active site loop is highly dynamic in both the inhibitor-free and CGS-27023A structures based on S² order-parameters (Moy, *et al.*, <u>J. Biomol. NMR</u> 1997). This region in the MMP-13:Compound A structure appears to be significantly less mobile by the observation that most of the residues in this loop region were easily observable in the ¹H-¹⁵N HSQC spectra and readily assigned. One possible explanation for this difference is the hydrophobic interaction between the pyridine ring of Compound A and the side-chain for Ile-140. In MMP-1, I140 is replaced by a serine which essentially eliminates this beneficial interaction.

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Another unique feature of the MMP-13 NMR structure is the apparent dynamic nature of residues H69 to Y73. These residues are completely disordered due to the lack of any assignment information and the resulting absence of any constraint information presumably a result of the flexible nature of these residues. Residues H69 to Y73 occur between the Ca⁺² binding loop and the structural zinc where the corresponding region in the previously solved MMP-1 NMR structures is well defined. There is no apparent explanation for this change in mobility between the two NMR structures but it may contribute to the observed difference in the physical behavior of MMP-1 and MMP-13. Under identical conditions, inhibitor-free MMP-1 is stable for upwards of two months whereas inhibitor-free MMP-13 degrades immediately.

Comparison of the MMP-13:Compound A and MMP-1:CGS-27023A Structures:

The high-resolution NMR structure for the MMP-13:Compound A complex was effectively and efficiently determined by using a homology model based on the

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MMP-1 NMR structure as an initial structure to analyze ambiguous NOESY data. This result is evident of the high structural and sequence similarity between members of the MMP family and consistent with the previously observed best-fit superposition of the backbone atoms for MMP-1, stromelysin, matrilysin and neutrophil collagenase (Moy, *et al.*, <u>Biochemistry</u> 1998; Moy, *et al.*, <u>Biochemistry</u> 1999).

The strong similarity between the various MMP structures creates an initial difficulty in designing specific MMP inhibitors. This is exemplified by the high sequence similarity among the MMPs in the active site. Comparison of the sequence similarity between MMP-13 and MMP-1 illustrates this difficulty. There are only a few significant residue differences between the two enzymes where these modifications results in a significant change in the local environment of the active site. The R114 to V115 modification results in a conversion from a hydrophilic to a hydrophobic environment at the base of the S1' pocket between MMP-1 and MMP-13, respectively. Similarly, the N80 to L81 substitution places a bulkier hydrophobic residue in the S2' pocket for MMP-13 compared to a more hydrophilic environment for MMP-1. Similarly in the active loop region, I140 a bulky hydrophobic residue in MMP-13 replaces the smaller hydrophilic S139 residue in MMP-1. Clearly, it is feasible to incorporate substituents into a small molecule to take advantage of these spatial distinct environmental changes between MMP-1 and MMP-13. Nevertheless, when these sequence and environmental differences are averaged across the MMP family it becomes less discriminating and extremely difficult to design an inhibitor to a specific MMP subtype based strictly on the small sequence differences.

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Conversely, the most distinct structural difference between the MMPs and readily amenable to incorporating specificity in drug design is the relative size and shape of the S1' pocket. This is clearly evident by comparison of the defined S1' pockets for MMP-13 and MMP-1. The large difference in size in the S1' pockets between the MMP-13 and MMP-1 NMR structures is striking. The S1' pocket for MMP-13 nearly reaches the outer surface of the protein and

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is greater then twice the size of MMP-1. The additional size of the MMP-13 S1' pocket relative to MMP-1 is best illustrated by the filling capacity of the two inhibitors. In the MMP-1:CGS-27023A NMR structure, the p-methoxyphenyl effectively fills the available S1' pocket for MMP-1. Conversely, in the MMP-13:Compound A complex the p-methoxyphenyl only partially fills the available space within the MMP-13 S1' pocket. The size of the MMP-13 pocket is actually similar in size to stromelysin where the design of stromelysin inhibitors has taken advantage of this deeper S1' pocket by using a biphenyl substituent in another series instead of the p-methoxyphenyl in Compound A to bind into the S1' pocket (Hajduk, et al., J. Am. Chem. Soc. 1997; Olejniczak, et al., J. Am. 10 Chem. Soc. 1997). Thus, the NMR structures for MMP-13 and MMP-1 suggest that a ready approach to designing specificity between these MMPs is to take advantage of the significantly different sized S1' pockets. The high mobility of the MMP-1 active site presents a potential caveat to this analysis of the static images of the MMP-1 and MMP-13 structures. It is probable that the MMP-1 active site is capable of accommodating a S1' substituent larger then implied from its current structure due to its increased mobility in both free and inhibited structures.

Examination of the binding mode of Compound A in the MMP13:Compound A complex suggests a conformation generally similar to CGS27023A in the MMP-1:CGS-27023A NMR structure previously reported (30 simulated annealing structures deposited with Protein Data Bank, Accession No.
4AYK; restrained minimized mean structure deposited with Protein Data Bank, Accession No. 3AYK). Compound A and CGS-27023A are structurally very similar with the only difference being the nature of the substituent binding in the S2' pocket where an aryl group in Compound A replaces the isopropyl group in CGS-27023A. The strong resemblance between the binding mode of Compound A and CGS-27023A is apparent from the nearly identical intermolecular NOE patterns observed between the inhibitors and the proteins.

The key MMP-13 residues involved in the interaction with Compound A correspond to L81, L82 and A83 from β-strand IV; residues L115, V116, and

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H119 from α -helix II; and L136, I140 and Y141 from the active site loop. Similarly, the MMP-1 residues involved in the interaction with CGS-27023A correspond to residues N80, L81, A82 and H83 from β -strand IV; residues R114, V115, H118 and E119 from α -helix II; and L135, P138, Y137, S139 and Y140 from the dynamic flexible loop.

As stated previously, there are three distinct residue changes between MMP-13 and MMP-1 in the active site. The R114 to L115 change between MMP-1 and MMP-13, respectively, has a significant impact on the environment at the base of the S1' pocket but since Compound A only partially fills the MMP-13 S1' pocket this change should not effect the binding conformation of Compound A relative to CGS-27023A. Conversely, the N80 to L81 substitution directly interacts with the inhibitors in the S2' pocket and may result in an effective change in the binding mode of the inhibitors. To complicate the analysis, the only change in the inhibitors are the substituents that bind the S2' pocket. For the MMP-1:CGS-27023A complex, the isopropyl 15 group interacts with both the sidechains of N80 and H83 where the aryl group from Compound A only interacts with L81 in MMP-13. Additionally, CGS-27023A is in hydrogen-bonding distance to both L81 and A82, whereas Compound A appears to form a bifurcated hydrogen bond with L82. This analysis suggests that CGS-27023A binds closer to β -strand IV since the S2' 20 pocket is more accessible in MMP-1 due to the absence of the bulky L81 sidechain and the presence of the aryl group in Compound A. A direct comparison of the bound conformations suggest only a subtle difference in the relative orientation of the inhibitors. The S139 to I140 difference between MMP-1 and MMP-13, respectively, appears to be related to a mobility change as opposed to a structural change. In the MMP-1:CGS-27023A structure the pyridine ring position is essentially undefined and solvent exposed this compares to the MMP-13:Compound A structure where the pyridine ring binds with the side-chain of I140. Clearly, Ile is a bulkier more hydrophobic group relative to Ser which would provide a beneficial hydrophobic interactions with the pyridine ring. The 30 more interesting observation is the apparent decrease in mobility for the active

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loop in the MMP-13 structure which may be related the pyridine ring I140 interaction. This appears to be consistent with previously inhibited MMP X-ray structures (Spurlino, et al., Proteins: Struct., Funct., Genet. 1994) where the inhibitor may extend the formation of a β -sheet between b-strand IV and the active loop region which results in low B-factors in the X-ray structure. This may suggest that the mobility of the active loop region is easily removed with any positive interaction with the inhibitor.

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There are apparently some interesting differences between the mode of binding for the two inhibitors in the MMP-13:Compound A and MMP-1:CGS-27023A NMR structures. The more striking observation is the overall similarity between the two structures. Despite some significant sequence differences and a large difference in the size and shape of the S1' pocket either inhibitor structure would accurately predict the other structure. This observation seems to indicate that the major contributing factors to inhibitors binding the MMPs is the fit in the S1' pocket and the binding of the hydroxamic acid to the catalytic zinc. The interaction in the S2' pocket appears to have a more subtle impact on inhibitor binding and selectivity since both Compound A and CGS-27023A are low nanomolar inhibitors of MMP-13 and MMP-1, respectively. Therefore, the high-resolution solution structure of the MMP-13:Compound A in conjunction with the previously reported MMP-1 NMR structures suggest that taking advantage of the significant differences in the size and shape of the S1' pocket is a reasonable approach for developing specific MMP inhibitors.

structure of MMP-13 complexed with a sulfonamide derivative of a hydroxamic acid compound (Compound A). The overall fold of MMP-13 is similar to previously reported MMPs structures. The major difference is the large S1' pocket which nearly reaches the surface of the protein. The structure was based on a total of 3279 constraints including 47 distance restraints between MMP-13 and Compound A from X-filtered NOESY experiments. The inhibitor was found to bind to the "right" side of the catalytic Zn such that the p-methoxyphenyl ring

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sits in the S1' pocket, the aryl moiety interacts with L81 of β IV, the pyridine ring interacts with I140 of the active site loop, hydrogen bond interactions exist between the sulfonamide oxygens with residue L82 and the hydroxamic acid chelates the catalytic Zn. This inhibitor binds MMP-13 similarly to the MMP-1: CGS-27023A complex suggesting that appropriately filling the S1' pocket may play a key role in developing selective MMP inhibitors.

Table 1. Observed NOEs Between Compound A and MMP-13

| Compound A | MMP-13 | NOE Class | Compound A | MMP-1 | NOE Class |
|-------------|-------------|-----------|------------|-----------|-----------|
| 1HH* | L81 Ηγ | W | 3HH* | Υ141 Ηα | M |
| 1HH* | L81 Hδ1# | W | 3HH* | Υ141 Нβ1 | W |
| 1HH* | L81 Hδ2# | M | 3HH* | Υ141 Ηβ2 | W |
| 1HH* | -L81 Hα | S | 3HH* | Υ141 Ηδ2 | W |
| 1HE2 | L81 H81# | W | 3HE2 | L82 Hδ1# | W |
| 1HE2 | L81 Hδ2# | M | 3HE1 | Α83 Ηβ# | w · |
| 1 HZ | L81 Hδ1# | W | 3HE1 | Н116 На | W |
| lHZ | L81 Hδ2# | M | 3HE1 | Η116 Ηγ1# | M |
| 2HZ | I140 Hγ2# | W | 3HE2 | Η116 Ηγ2# | W |
| 2HE1 . | I140 Ηδ1# | W | 3HE2 | I140 Ηγ2# | W |
| 3HH* | L82 Hδ1# | W | 3HE2 | Υ141 Ηα | W |
| 3HH* | L115 Hβ# | W | 3HE2 | Ύ141 Ηβ1 | W |
| 3HH* | L115 Hy | W | 3HE2 | Y141 Hβ2 | W |
| 3HH* | L115 Hδ1# | W | 3HD2 | L82 H81# | W |
| 3HH* | - L115 Hδ2# | W | 3HD1 | Α83 Ηβ# | W |
| 3HH* | V116 Ha | w | 3HD1 | V116 Hy1# | W |
| 3HH* | V116 Hy1# | W | 3HD2 | V116 Hγ2# | W |
| 3HH* | V116 Hy2# | M | 3HD2 | 1140 Ηα | W |
| 3HH* | Η119 Ηα | W | 3HD2 | I140 Ηγ2# | W |
| 3HH* | Н119 Нδ2 | W | 3HD2 | Υ141 Ηα | W |
| 3HH* | Н119 НВ1 | W | 3HD2 | Υ141 Ηβ1 | W |
| 3HH* | Н119 Нβ2 | w | 3HD2 | Υ141 Ηβ2 | W |
| 3HH* | L136 Hδ1# | W | 3HD2 | Y141 HN | W |
| 3HH* | L136 Hδ2# | W | | | |

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Example 3

Structure Based Design of a Novel, Potent, and Selective Inhibitor for MMP-13

The matrix metalloproteinases (MMPs) comprise a family of zinc containing enzymes that cleave a broad range of substrates including collagens, fibronectin and gelatins where the substrate preference various for individual MMPs. The design of MMP inhibitors has been initially based upon imitation of the binding interaction of natural protein substrates to MMPs where structural information of MMPs complexed with peptide substrates has been determined by x-ray crystallography and NMR spectroscopy. This structural information has provided a general description of the MMPs active site.

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The active site for the MMPs is composed of a catalytic zinc chelated by three histidines where three substrate binding pockets are located to both the right (S1', S2', S3') and left (S1, S2, S3) of the catalytic zinc. The substrate binding pockets were identified by the interactions of side chains from the peptide substrate with the MMPs. The primary effort in MMP inhibitor design has focused on compounds that chelate the catalytic zinc while primarily binding in the S1' and S2' pockets. This has evolved from the observation that the structural characteristics of the S1' pocket (size, shape, amino acid composition) incurs the greatest variability between the individual MMPs and this provides an obvious approach in designing selective and specific MMP inhibitors. Nevertheless, there has also been success in utilizing the binding pockets to the left of the catalytic zinc in addition to or in combination with the right handed binding pockets in the design of inhibitors.

The underlying challenge in designing MMP inhibitors is the reasonably high sequence and structural homology observed between the individual members of the MMP family making it intrinsically difficult to design an inhibitor that will function against a single MMP in the absence of structural information. The problem with a non-specific MMP inhibitor as a drug is the high likelihood of serious side-effects because of the large number of enzymes in the MMP family and their corresponding diversity in targets and function.

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Accordingly, the detailed structural information provided herein is a critical component of an inhibitor design program targeting a particular MMP enzyme.

Materials and Methods:

5 Synthesis of Compound D and Compound E: The sulfonamide derived from 2-amino-3,5-dimethyl-benzoic acid methyl ester and p-methoxybenzenesulfonyl chloride was N-alkylated with benzyl bromide and the ester group of the resulting intermediate was hydrolyzed (LiOH/THF) to afford the carboxylic acid. The corresponding hydroxamic acid was formed by preparation of the acid chloride (oxalyl chloride/DMF) followed by reaction with hydroxylamine. Compound E was synthesized by reaction of 2-amino-3,5-dimethyl-benzoic acid methyl ester and p-fluorobenzenesulfonyl chloride followed by N-alkylation with benzyl bromide. Hydrolysis of the methyl ester (LiOH/THF) followed by displacement of fluorine with the alkoxide of benzofuran-2-carboxylic acid (2-hydroxy-ethyl)-amide gave, after conversion to the hydroxamic acid and formation of the HCl salt as described above, Compound E.

NMR Sample Preparation: Uniformly (>95%) ¹⁵N- and ¹⁵N/¹³C-labeled human recombinant MMP-13 was expressed in E. coli and purified as described previously. 1mM ¹³C/¹⁵N- and ¹⁵N- MMP-13 NMR samples were prepared by concentration and buffer exchange using Millipore Ultrafree -10 centrifugal filters into a buffer containing 10mM deuterated Tris-base, 100mM NaCl, 5mM CaCl₂, 0.1 mM ZnCl₂, 2 mM NaN₃, 10mM deuterated DTT in 90% H₂O/10% D₂O or 100% D2O. The 10:1 Compound B:MMP-13 samples were prepared by addition of Compound B into either a 1mM ¹³C/¹⁵N- or ¹⁵N-MMP-13 sample 25 followed by pH readjustment. The sample to explore the potential of competitive inhibition between Compound B and Compound A was prepared by first adding 1mM of Compound A to a 1 mM ¹⁵N- MMP-13 sample followed by the addition of 10mM Compound B. The initial MMP-13:Compound A sample was made by buffer exchange of ¹⁵N- MMP-13 into the buffer containing 0.1 30 mM Compound A followed by additional buffer exchanges to remove excess

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Compound A. Finally, 10mM of Compound B was added to the 1mM ¹⁵N- MMP-13:Compound A sample followed by pH readjustment.

NMR Data Collection: All spectra were recorded at 35°C on a Bruker AMX-2 600 spectrometer using a gradient enhanced triple-resonance ¹H/¹³C/¹⁵N probe. For spectra recorded in H₂O, water suppression was achieved with the WATERGATE sequence and water-flip back pulses (Piotto, et al., J. Biomol. NMR 1992; Grzesiek and Bax, J. Am. Chem. Soc. 1993). Quadrature detection in the indirectly detected dimensions were recorded with States-TPPI hypercomplex phase increment (Marion, et al., J. Magn. Reson. 1989). Spectra were collected with appropriate refocusing delays to allow for 0,0 or -90,180 phase correction.

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The resonance assignments and bound conformation of Compound A in the MMP-1: Compound A complex were based on the 2D ¹²C/¹²C-filtered NOESY (Petros, *et al.*, <u>FEBS Lett.</u> 1992; Gemmecker, *et al.*, <u>J. Magn. Reson.</u> 1992), 2D ¹²C/¹²C-filtered TOCSY (Petros, *et al.*, <u>FEBS Lett.</u> 1992; Gemmecker, *et al.*, <u>J. Magn. Reson.</u> 1992) and ¹²C/¹²C-filtered COSY experiments (Ikura and Bax, <u>J. Am. Chem. Soc.</u> 1992).

The assignments of the ¹H, ¹⁵N, and ¹³C resonances of MMP-13 in the MMP-13:Compound B complex were based on the previous assignments for the MMP-13:Compound A complex in combination with a minimal set of experiments: 2D ¹H-¹⁵N HSQC, 3D ¹⁵N- edited NOESY (Marion, *et al.*<u>Biochemistry</u> 1989; Zuiderweg and Fesik, <u>Biochemistry</u> 1989), CBCA(CO)NH (Grzesiek and Bax, <u>J. Am. Chem. Soc.</u> 1992), C(CO)NH (Grzesiek, *et al.*, <u>J. Magn. Reson.</u>, Ser. <u>B</u> 1993), HNHA (Vuister and Bax, <u>J. Am. Chem. Soc.</u> 1993) and HNCA (Kay, *et al.*, <u>J. Magn. Reson.</u> 1990). The acquisition parameters for each of the experiments used in determining the solution structure of the MMP-13:Compound B complex were as reported previously (Moy, *et al.*, <u>Biochemistry</u> 1996).

The MMP-13:Compound B structure is based on observed NOEs from the 3D ¹⁵N-edited NOESY (Marion, et al. <u>Biochemistry</u> 1989; Zuiderweg and Fesik, <u>Biochemistry</u> 1989) and 3D ¹³C-edited/¹²C-filtered NOESY (Vuister

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and Bax, <u>J. Am. Chem. Soc.</u> 1993; Lee, et al., <u>FEBS Lett.</u> 1994). The 3D ¹⁵N-edited NOESY and 3D ¹³C-edited/¹²C-filtered NOESY experiments were collected with 100 msec and 110 msec mixing times, respectively.

Molecular Analysis and Design: The minimized models of Compound B and Compound D complexed to MMP-13 were prepared as previously described (Chen, et al., J. Biomol. Struct. Dyn. 1995; Chen, et al., Biochemistry (in press) 1998). Using molecular dynamics methods (Sybyl v6.4 from Tripos Inc), protein regions within 5 Å from Compound B were sampled along with the inhibitor, 10 whereas everything else remained rigid during the simulations. Upon energy convergence, the last 50 frames from the final 100 picoseconds run was averaged and this averaged structure underwent a final minimization. The final protein-Compound B model appeared to have optimized possible polar and van der waals interactions. The identical procedure was applied to the complex of MMP-13 and Compound D. Since the two complexes used identical MMP-13 15 structures, the proteins were overlapped to depict the positions of the two inhibitors within the active site. Graphics analysis of the inhibitors showed that the methylene carbon of Compound B containing the 2HB1/2 protons (Figure 6) overlapped identically with the methoxy carbon from Compound D. This analysis indicated the optimal or minimal linkage length of connecting the 20 benzofuran moiety to the methoxy region of Compound D. The final design scheme is shown in Figure 8A for the hybrid inhibitor. The homology model of MMP-9 was constructed using the COMPOSER program (Tripos INC, Sybyl v.6.4)

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High-throughput Screening Analysis: Compound B was identified as an initial lead from the analysis of the MMP-13 high-throughput screen (HTS). A total of 58079 compounds were screened for their ability to inhibit MMP-13 enzymatic activity where 385 compounds were shown to have \geq 40% inhibition at 10 μ g/ml dosage. Compound B was shown to exhibit weak inhibition of MMP-13 (89% at the 10 μ g/ml), but more intriguing was the observation of a complete

lack of activity against other MMPs (MMP-1, MMP-9 and TACE). The primary structure of Compound B along with the proton naming convention is shown in Figure 6.

The resulting HTS hits were further examined by cluster analysis.

The hits were clustered based on structural similarities where the properties of these compounds were compared against the properties of the set of orally available drugs. The properties used to profile the HTS hits consists of: total number of non-hydrogen atoms, number of heteroatoms, number of hydrogen-bond donors and acceptors, calculated logP and molecular weight. This profile analysis provides an initial means to predict the likelihood that an HTS hit may have drug-like characteristics such as bioavailability and in-vivo stability. The profile of Compound B indicates that the compound has properties similar to orally available drugs suggesting that it would be an ideal candidate for optimization of its enzyme potency and selectivity.

A common feature of known MMP inhibitor structures is the presence of a Zn-chelator that plays a fundamental role in its activity. In most cases Zn chelation occurs from the presence of a hydroxamic acid in the structure of the small molecule. As apparent from the structure of Compound B, the compound does not contain an obvious substituent that would chelate Zn.

Thus, the unique structure of Compound B suggested a potential novel mechanism for inhibition of MMP-13 further strengthening the choice of Compound B as an initial lead candidate. Therefore, the identification of Compound B as a candidate to optimize its activity and selectivity was based on three unique observations: its intrinsic MMP-13 selectivity, its structural profile similar to known bioavailable drugs and finally its apparent novel structure.

NMR Structure of the MMP-13 - Compound B Complex: The NMR binding studies provided critical information pertaining to the mechanism of Compound B inhibition of MMP-13 and the method for designing increase potency. The major question presented when Compound B was identified from HTS was its unknown MMP-13 binding site and its method for inducing MMP-13 inhibition.

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Previous work on the NMR structure of MMP-13 complexed with Compound A and MMP-1 complexed with CGS-27023A provided the framework and methodology to analysis Compound B bound to MMP-13 (Moy, et al., Biochemistry Submitted 1999; Moy, et al., Biochemistry 1999).

The Compound B MMP-13 binding site was initially identified 5 from chemical shift perturbation in the ¹H-¹⁵N HSQC spectra. The observed perturbations were mapped onto a GRASP surface (not shown). It is apparent that the major effect of Compound B on the chemical shifts of MMP-13 occurs in the proximity of the S1' pocket suggesting that Compound B sits in this pocket. From the NMR and X-ray structures of MMP-13, it was determined that the S1' 10 pocket for MMP-13 is very deep and linear in shape while nearly reaching the surface of the protein. In fact, a number of residues at the surface of MMP-13 near the base of the S1' pocket show significant chemical shift perturbation in the presence of Compound B. Since Compound B is a linear molecule, docking studies would place the inhibitor stretched throughout the linear S1' pocket of MMP-13. The only question remaining was whether to place the morpholine or the benzofuran moiety of Compound B at one end of the pocket, adjacent to the catalytic zinc or the opposite end, distant from the zinc atom. Property analysis of the enzymes S1' pocket depicts that the end adjacent to the zinc is relatively polar whereas the opposite end is hydrophobic. This analysis lead us to dock 20 Compound B with the morpholine ring adjacent to the catalytic zinc atom with the benzofuran moiety siting in a hydrophobic pocket formed by L115, L136, F149 and P152 at the base of the S1' pocket. To further verify the proposed binding of Compound B in the S1' pocket of MMP-13, a simple competition experiment with Compound A was conducted. The ¹H-¹⁵N HSQC experiment for 25 the MMP-13:Compound B complex was collected in the presence of Compound A. The presence of Compound A displaced all of Compound B as evident by the distinct differences in the ¹H-¹⁵N HSQC spectra which further suggests that both compounds bind in the S1' pocket.

The relative orientation and binding of Compound B with MMP-13 was further confirmed by the observation of intermolecular NOEs between

Compound B and MMP-13 from the 3D ¹³C-edited/¹²C-filtered NOESY experiment. The NOESY spectra was collected in the presence of a ten-fold excess of Compound B because of the weak affinity of Compound B with MMP-13. Nevertheless, a total of 16 NOEs were observed between Compound B and L81, L115, V116, Y141, T142 and Y143 which support the initial positioning of Compound B in the MMP-13 S1' pocket. An expanded 2D plane from the 3D ¹³C-edited/¹²C-filtered NOESY experiment (not shown) demonstrated examples of some key intermolecular NOEs between Compound B benzofuran group resonances and L115 δ and Compound B resonances proximal to the morpholine ring and L82 δ. The complex of Compound B with MMP-13 was 10 subjected to energy refinement using the NMR results as constraints (Moy, et al., Biochemistry 1999; Chen, et al., J. Biomol. Struct. Dyn. 1995). The modeling results depict the morpholine oxygen forming a hydrogen bond with the backbone amide group of Leu-82 and the benzofuran group packs deep in the S1' pocket with the peptide bond linker portion forming hydrogen bonds 15 with protein backbone groups. The complex shows no apparent interactions between the inhibitor and the catalytic zinc justifying the ligands micromolar potency.

20 Structures of MMP-1, MMP-9 and MMP-13: The recent NMR solution structures of MMP-1 and MMP-13 were used as starting points for molecular modeling and analysis (Moy, et al., Biochemistry Submitted 1999; Moy, et al., Biochemistry 1998; Moy, et al., Biochemistry 1999). A homology model for MMP-9 was developed based on its strong homology to MMP-1 (54% identity around the catalytic domain). Based on the homology model, the catalytic site of MMP-9 is similar to the corresponding sites in MMP-1 and MMP-13. All three structures were used as starting points for analysis and synthetic design.

Comparative analysis of the MMP structures shows that residue positions 115 and 144, in addition to the length of the specificity loop, determines the size and shape of the S1' pockets. Alignment of the NMR structures for MMP-1 and MMP-13 shows that MMP-13 contains two additional

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insertions in the specificity loop. The homology model of MMP-9 indicates no additional insertions so its length is identical to MMP-1.

Residue positions 115 and 144 are important in establishing the relative length of the S1' pockets for the MMPs where the larger the side chain at these positions results in a smaller S1' pocket. Since residue 115 is spatially closer to the catalytic zinc than residue 144, a larger side chain for residue 115 will have a greater impact on defining a smaller S1' pocket compared to residue 144. MMP-1 has the largest side chain at position 115, thus its S1' pocket is the smallest. MMP-9 has an Arg at position 144 resulting in its S1' pocket being longer compared to MMP-1. Conversely, MMP-13 has short side chains at both positions 115 and 144. The short side chains combined with an increased length of its specificity loop result in MMP-13 having the largest S1' pocket. To summarize, the size of the MMP S1' pockets are as follows: MMP-13 > MMP-9 > MMP-1 where this structural feature plays a critical role in the design strategy for developing a potent and specific MMP-13 inhibitor.

Design Strategy: A strategy utilizing NMR and molecular modeling was applied towards the design and synthesis of an MMP-13 selective inhibitor lead. The basic approach behind the design strategy is to optimize the affinity of the chemical lead Compound B while maintaining its inherent MMP-13 selectivity. This can be achieved by taking advantage of the distinct structural feature of MMP-13, its deep linear S1' pocket, while combining overlapping structural features of Compound B with other potent inhibitors. Compound C is an example of a potent and selective inhibitor for MMP-9 and MMP-13 (See Table 2). Based on the NMR solution structure of MMP-13 complexed with Compound A (Figure 4), structurally similar inhibitors were positioned into the active site of MMP-13.

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Figure 7 shows the critical regions of Compound C, which can be broken down into two components, Compound D which represents the zinc chelating portion of the compound that contributes to the binding potency and the toluene group (1A) which contributes to enhanced ligand selectivity against

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MMP-1. The strategy was to design a new inhibitor based on replacing the toluene group (1A) with a component of Compound B critical for binding within the extended S1' pocket of MMP-13. The overlay of the NMR solution structure for Compound B with the model for Compound D is shown in Figure 8B. The close similarity between the positioning of the two structures made it readily apparent that it would be possible to generate a hybrid of the two structures combining the potent Compound D with the selective component of Compound B (Figure 8A). These results were then used to design the proposed hybrid inhibitor Compound E. The assay data in Table 2 clearly shows that the new inhibitor, Compound E, has better potency compared to Compound C in addition to improved selectivity towards MMP-13. Thus, the combination of NMR spectroscopy with molecular modeling techniques resulted in the design of a novel, potent and selective MMP-13 inhibitor (Compound E) which has an IC50 of 17 nM for MMP-13 and showed >5800, 56 and >500 fold selectivity against MMP-1, MMP-9 and TACE, respectively. To the best of our knowledge, this represents the first example of a potent MMP-13 inhibitor that has been shown to be selective against MMP-9.

Table 2 - IC50 and Selectivity Data

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| Compoun d | MMP-1 | MMP-9 | MMP-13 | TACE | S-1ª | S-9ª | S-TACE [®] | | | |
|--------------------------|--|-------|--------|-------|--------|------|---------------------|--|--|--|
| С | 750nM | 46nM | 75nM | 470nM | 10.0x | 0.6x | 6.3x | | | |
| D | 82nM | 21nM | 15nM | 240nM | 5.5x | 1.4x | 16x | | | |
| Е | NA | 945nM | 17nM | 19% | >5800x | 56x | >500x | | | |
| F | 1025n M | 71nM | 301nM | 664nM | 3.4x | 0.2x | 2.2x | | | |
| ^a Selectivity | ^a Selectivity data presented as a ratio of the MMP or TACE IC50 with MMP-13 | | | | | | | | | |

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Example 4

The X-ray crystal structure of the MMP-13:Compound A complex was determined using the following procedure:

5 Gene/expression system/production: The cDNA coding for human MMP-13 proenzyme had 85 residues of the PRO domain, followed by 165 residues of the catalytic domain (CAT). The gene was carried on a pET-21a expression plasmid, under the control of a bacteriophage T7 promoter. The expression host was Escherichia coli BL21(DE3), which had a chromosomal copy of T7 RNA polymerase under lac control. Cells were grown in nutrient broth, and synthesis of PRO-CAT was induced by isopropyl-β-thiogalactoside. The protein accumulated to 5-10% of total cellular protein, essentially all of which was aggregated into inclusion bodies.

For potential MAD experiments, the plasmid was transferred into a methionine auxotroph host. PRO-CAT with selenomethionine substitution was produced by induction in a defined medium, with methionine replaced by selenomethionine.

Purification and refolding of PRO-CAT: Frozen cells were disrupted
mechanically, and inclusion bodies were isolated by centrifugation. PRO-CAT was solubilized with urea containing dithiothreitol to disrupt any disulfide bridges. PRO-CAT was partially purified by anion-exchange chromatography, in urea, on Q Sepharose. The protein was diluted to about 400 μg/ml in a solution of sodium chloride, calcium chloride, and zinc acetate, buffered with tricine-HCI. Refolding proceeded over 3-4 days, during dialysis, with multiple buffer changes. PRO-CAT was then concentrated for activation and release of CAT.

Activation of PRO-CAT: The presently-accepted view of MMPs holds that the proenzyme form is maintained in an inactive state through the coordination of one cysteine from the PRO domain into the active-site zinc. If this cysteine is

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displaced, the enzyme becomes active. In our protocol, aminophenyl mercuric acetate was added to the protein solution to form a mercurial adduct with the cysteine. Progress of activation was monitored by SDS polyacrylamide gel eletrophoresis. Results indicated that the CAT domain accumulated and the PRO domain was degraded to small peptides.

Purification of MMP-13 (CAT) – Size Exclusion: Following activation and PRO cleavage, MMP-13 was isolated by size-exclusion chromatography through SuperDex 75 in a solution of sodium chloride, calcium chloride, and zinc acetate, buffered with tris-HC1.

Purification of MMP-13 – Affinity: MMP-13 was further purified by affinity chromatography on an immobilized hydroxamate inhibitor. The affinity matrix was prepared by coupling an hydroxamate inhibitor to Sepharose through the amino group of the piperazine ring. MMP-13 can be absorbed to the matrix and desorbed by displacement using another inhibitor of choice.

Characterization of MMP-13: Protein preparations for crystallization trials were validated by several techniques. Routinely, SDS-PAGE showed a predominant species whose migration was consistent with a molecular weight of around 19,000. MALDITOF mass spectroscopy demonstrates a single species consistent with the expected size of 18,588 amu. (MMP-13 prepared with selenomethionine showed essentially complete replacement). N-terminal sequencing demonstrated that the protein begins with YNVF, as expected for correct cleavage between PRO and CAT. Retention volume in analytical size-exclusion chromatography was consistent with a monomeric protein: no detectable aggregation was observed. The final protein was enzymatically active on a fluorogenic peptide substrate, and degraded denatured collagen.

30 Crystallization of MMP-13 complex with Compound A: The MMP-13 protein solution was buffered with 10 mM tris-HCL buffer, pH 7.5, and 0.25 M NaCl.

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The concentration of protein used for crystallization was 20.0 mg/ml. The inhibitor solution was added to a protein solution with a mole ratio (protein:inhibitor) of 1:2, and was mixed for more than 1 hour.

Crystallization conditions were screened by the hanging-drop vapor diffusion method (Mcpherson, A., Methods Biochem. Anal. 1976). A successful procedure for growing crystals of this complex at room temperature was identified, and crystals were obtained. Specifically, a solution was prepared from 3 µl of protein solution and 3 µl of precipitant solution, which consisted of 26% PEG4000, 0.1 M ammonium sulfate, and 0.1 M sodium chloride. A drop of this solution was suspended on a microscope coverslip glass which had been 10 coated with silicone to prevent drop spreading. The reservoir solutions consisted of 0.6 ml precipitant solution. Equilibration was performed at room temperature by vapor diffusion. Crystals began appearing after three days. After two weeks, these crystals stopped growing. The X-ray data which have been processed show that the MMP-13 complex was crystallized in two forms. One crystal form is C-centered orthorhombic; it belonged to space group C2221, and had a cell dimension of a=36.3 Å, b=134.4 Å, and c=134.8 Å. This crystal had high mosaicity; therefore, it would be of little use when working on the structure of the complex. The second crystal form is primitive orthorhombic, from space group P21212, with a cell constant of a=108.3 Å, b=79.8 Å, and c=36.1 Å. This crystal had low mosaicity, but it was very small in most cases.

In order to obtain a big single crystal for X-ray data collections, the seeding technique (Thaller, C., et al., J. Mol. Biol. 1981) was applied. This was accomplished by using both the microseeding and the macroseeding methods. Small seed crystals were transferred to a 20% PEG4000 precipitant solution on a depression slide. A single washed crystal was injected into a hanging-drop solution, which was composed of 3 μ l of MMP-13 complex solution and 3 μ l of precipitant solution. The reservoir solutions consisted of 0.6 ml precipitant solution at pH 8.0. This procedure successfully produced bigger crystals with a maximum edge dimension of up to 0.35 x 0.1 x 0.1 mm³. These crystals diffracted X-ray at a resolution of 2.0 Å.

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X-Ray Data Collection: X-ray diffraction data from 30.0-2.0 Å resolution for the MMP-13:Compound A complex crystal (P21212 form) was collected by using an RAXIS IIc Image Plate area detector which used graphite monochromatic CuKα radiation from a Rigaku RU200 rotating anode generator (operating at 50 kV, 100 mA) at a low temperature of 100 K. The oscillation angle for each plate was 1 degree, and exposure time was 20 minutes per 'image'. The processing of X-ray diffraction data was accomplished using the HKL programs (Otwinowski, Z. and Minor, W., Methods in Enzymology 276:307-26). The R-merges for full and partial reflections were 4.0% and 6.04% respectively. 18,782 unique
reflections (81% complete at 2.0 Å resolutions) were collected.

Structure Determination and Refinement: The MMP-13 complex crystal structure has been determined by a combination of crystallographic modeling and the Molecular Replacement method using models of MMP-13 derived from the MMP-1 and MMP-8 structures. The homology between MMP-13 and MMP-8 is 56% by sequence, and at least 70% by structure. Crystals of the MMP-13 complex have two molecules in the asymmetric unit, *i.e.*, the unit is a dimer. Conventional molecular replacement was not effective for determination of this dimer structure by using a monomer model. There are two reasons for this: (1) the high symmetry of the crystal structure; and (2) the conformations and the configurations of the side chain and the main chain in flexible loops of MMP-13 and MMP-8.

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Firstly, the crystal structure of the MMP-13 complex is highly symmetrical. The P21212 crystal has four symmetry operations, and there are eight molecules in a unit cell. A second crystal form, belonging to space group C222, and having eight symmetry operations in a unit cell, has been identified. In this crystal, there are 16 monomers per cell in the dimer structure, and 32 monomers per cell in the tetramer structure. Therefore, the rotation search and especial translation search become more difficult. Secondly, even though the MMP family's catalytic domain structure is highly conserved, the conformations and the configurations of the side chain and the main chain in flexible loops of

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MMP-13 and MMP-8 may not be the same. In particular, the similarity between the two structures may not be sufficient to permit the determination of the dimer structure using a monomer as the searching model.

Many attempts at a rotation and translation search were made by using the X-ray data and models of either a monomer of MMP-8 or a dimer of MMP-1. Some rotation solutions were obtained, but no final translation solution has been found by using the monomer model. Accordingly, to determine this structure, it was proposed that a dimer model be constructed first; the molecular replacement method was then applied to solve the structure.

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The key idea of this proposal was crystal packing. To construct a dimer, the orientations of each monomer were determined on the basis of a rotation search. The positions of each monomer were located on the basis of the molecular packing in unit cell. Many dimer models have been constructed and applied as the 'model' for searching the rotation and translation using program AMORE (Collaborative Computational Project, Number 4 (CCP4) (1994), Acta Cryst. D50:760-763). One dimer model was found to be correct, and finally resulted in the MMP-13 3-D crystal structure using the molecular replacement method. The MMP-13 complex structure was confirmed by observing the most important and significant fact that the positions of the two zinc ions and the two calcium ions could be identified from the difference (Fo-Fc) maps with five-sigma cut, where Fo was observed structure factor and Fc was the calculated structure factor of the dimer model without zinc and calcium atoms.

These ions were located in the exact positions where they were

observed in other MMP family members. The molecule fits the (2Fo-Fc)
electron densities very well, both in main chain and in side chain. The molecule
fits the 2Fo-Fc electron density quite well. All of these MMP molecules are
conserved in the core structure region, especially the position of the central
helix and the catalytic zinc. The MMP-13 dimer structure was further confirmed
by applying the molecular replacement programs XPLOR (Brünger, A.T., XPLOR
Version 3.1 Manual, Yale University, New Haven CT) and MERLOT (Fitzgerald,

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P., MERLOT, version 2.4 (Nov. 10, 1991). All of them worked very well, and produced results which were in agreement with the MMP-13 structure.

Structure Refinement: The structure refinement was carried out by the program XPLOR. The initial dimer model included 320 amino acid residues without zinc and calcium ions. The dimer model was refined against 2.0 Å X-ray data, collected on an RAXIS IIc area detector at a temperature of 100 K. The progress of the refinement was evaluated from the quality of the protein molecular conformations and the electron density maps, and the values of the crystallographic R-factor. The initial R-factor was 52%. After rigid-body 10 minimization, conjugated-gradient minimization, a heating stage, a slow-cooling stage in the range from 4000K to 300K, energy minimization, B-factor refinement, and positional refinement, the R-factor lowered to 0.32. Electrondensity maps with coefficients of (2Fo-Fc) and (Fo-Fc), as well as the phases, were calculated. The difference map shows four zinc ions and four calcium ions in the dimer structure with five-sigma cut. Some side chain loops and a few main loops were rebuilt on the interactive graphics system. The rebuilt dimer plus the zinc and calcium ions, as the new model, was refined. The R-factor was down to 26.6%. At this stage, a model of inhibitor Compound A was positioned in the active-site region based on the difference electron density.

The complex structure was refined by repeating the above steps, with the R-factor down to 20%. The water molecules were modeled as oxygen atoms. Their initial positions were located by searching the peaks in the (Fo-Fc) difference maps. These positions were then checked by calculating the distance between 'water' and the oxygen and nitrogen of the protein. Together with the protein (complex) atoms, these 'water' molecules were refined against the X-ray data. Once the temperature factor of water was higher than 50, this water was omitted. 120 water molecules near the protein were found, and five water molecules were identified in the active site of each monomer. The (2Fo-Fc) maps were used to adjust the solvent model and to aid in the placement of new solvent molecules, as well as to check and correct the whole model. The r.m.s.

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deviations of $C\alpha$ atoms for bond angles and bond distances from ideal geometry were 1.6° and 0.012 Å. The final crystallographic R-factor was 22%, at a resolution of 2.0 Å.

All publications mentioned herein above, whether to issued

5 patents, pending applications, published articles, protein structure deposits, or
otherwise, are hereby incorporated by reference in their entirety. While the
foregoing invention has been described in some detail for purposes of clarity
and understanding, it will be appreciated by one skilled in the art from a
reading of the disclosure that various changes in form and detail can be made

10 without departing from the true scope of the invention in the appended claims.

What is claimed is:

A solution comprising a biologically active catalytic 1. fragment of human collagenase-3 (MMP-13) complexed with N-Hydroxy-2-[(4methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide ("Compound A").

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- 2. The solution of Claim 1, wherein the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1.
- 3. The solution of Claim 2, comprising 1 mM MMP-13 complexed with Compound A in a 1:1 molar ratio, in a buffer comprising 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl₂, 0.1mM ZnCl₂, 2mM NaN₃, and 10 mM deuterated DTT in either 90% H₂O/10% D₂O or 100% D₂O.
- The solution of Claim 3, wherein the MMP-13 is either ¹⁵N 4. enriched or ¹⁵N, ¹³C enriched.
- The solution of Claim 1, wherein the secondary structure of 5. the catalytic fragment of MMP-13 comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands.
- 6. The solution of Claim 5, wherein the alpha helices and beta strands are configured in the order β_{I} , α_{A} , β_{II} , β_{III} , β_{IV} , β_{V} , α_{B} , and α_{C} .
- The solution of Claim 6, wherein the three alpha helices 7. correspond to residues 28-44 (α_A), 112-123 (α_B) and 153-163 (α_C) of Figure 1, and the five beta strands correspond to residues 83-86 (β_{I}), 95-100 (β_{II}), 59-66 (β_{III}) , 14-20 (β_{IV}) , and 49-53 (β_{V}) of Figure 1.
- 8. A crystallized catalytic fragment of MMP-13 complexed with N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-

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methyl-benzamide ("Compound A").

- 9. The crystallized complex of Claim 8, wherein the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1.
- 10. The crystallized complex of Claim 9, characterized as being in orthorhombic form with space group P21212, and having unit cell parameters of $a=108.3\text{\AA}$, $b=79.8\text{\AA}$, and $c=36.1\text{\AA}$.
- 11. The crystallized complex of Claim 10, further characterized as consisting of two molecules of MMP-13:Compound A complex in the asymmetric unit.
- 12. The crystallized complex of Claim 11, wherein the secondary structure of the catalytic fragment of MMP-13 comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands.
- 13. The crystallized complex of Claim 12, wherein the alpha helices and beta strands are configured in the order β_{I} , α_{A} , β_{II} , β_{III} , β_{IV} , β_{V} , α_{B} , and α_{C} .
- 14. The crystallized complex of Claim 13, wherein the three alpha helices correspond to residues 28-44 (α_A), 112-123 (α_B) and 153-163 (α_C) of Figure 1, and the five beta strands correspond to residues 83-86 (β_I), 95-100 (β_{II}), 59-66 (β_{III}), 14-20 (β_{IV}), and 49-53 (β_V) of Figure 1.
- 15. An active site of MMP-13, characterized by a catalytic zinc, a beta strand, a Ca²⁺ binding loop, an alpha helix, and a random coil region.

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- 16. The active site of Claim 15, wherein the beta strand comprises residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1, the Ca²⁺ binding loop comprises residues F75, D76, G77, P78, and S79 according to Figure 1, the alpha helix comprises residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1, and the random coil region comprises residues P139, I140, and Y141 according to Figure 1.
- 17. The active site of Claim 16, wherein said active site comprises the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 18. The active site of Claim 17, further comprising the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 19. The active site of Claim 18, further comprising the relative structural coordinates of amino acid residues F149 and P152 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å.

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- 20. An active site of MMP-13 comprising the relative structural coordinates of a catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, \pm a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 21. A method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of:
- (a) using a three dimensional structure of MMP-13 as defined by the relative structural coordinates of amino acids encoding MMP-13 according to Figures 4 or 5, \pm a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å;
- (b) employing said three-dimensional structure to design or select a potential inhibitor or activator; and
- (c) synthesizing or obtaining said potential inhibitor or activator.
- 22. The method according to Claim 21, wherein the potential inhibitor is designed de novo.
- 23. The method according to Claim 21, wherein the potential inhibitor is designed from a known inhibitor.
- 24. The method of Claim 22, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 25. The method of Claim 23, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.

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- 26. The method according to Claim 21, wherein the step of employing the three dimensional structure to design or select the potential inhibitor comprises the steps of:
- (a) identifying chemical entities or fragments capable of associating with MMP-13; and
- (b) assembling the identified chemical entities or fragments into a single molecule to provide the structure of the potential inhibitor.
- 27. The method according to Claim 26, wherein the potential inhibitor is designed de novo.
- 28. The method according to Claim 26, wherein the potential inhibitor is designed from a known inhibitor.
- 29. The method of Claim 27, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 30. The method of Claim 28, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 31. An inhibitor identified or designed by the method of Claim 21.
- 32. An inhibitor identified or designed by the method of Claim 26.

| YNVFP | RTLKW | SKMNL | TYRIV | NYTPD |
|--------------|--------------|-------------|-------|-------|
| 5 | 10 | 15 | 20 | 25 |
| MTHSE | VEKAF | KKAFK | VWSDV | TPLNF |
| 30 | 35 | 40 | 45 | 50 |
| TRLHD | GIADI | MISFG | IKEHG | DFYPF |
| 55 | 60 | 65 | 70 | 75 |
| DGPSG | LLAHA | FPPGP | NYGGD | AHFDD |
| 80 | 85 | 90 | 95 | 100 |
| DETWT | SSSKG | YNLFL | VAAHE | FGHSL |
| 105 | 110 | 115 | 120 | 125 |
| GLDHŞ | KDPGA | LMFPI | YTYTG | KSHFM |
| 130 | 135 | 140 | 145 | 150 |
| LPDDD 155 | VQGIQ 160 | SLYG 164 | - | |

FIG. 1

Sequence 1: MMP-13 Sequence 2: MMP-1

Identity score:

58.9 %

VGEYNVFPRTLKWSKMNLTYRIVNYTPDMTHSEVEKAFKKAFKVWSDVTPLNFTRLHDGIADIMISFGIKEHGDFYPFDG LTEGN PR WEQTHLTYRIENYTPDLPRADVDHAIEKAFQLWSNVTPLTFTKVSEGQADIMISFVRGDHRDNSPFDG

PSGLLAHAFPPGPNYGGDAHFDDDETWTS

SSKGYNLF

LVAAHEFGHSLGLDHSKDPGALMF

PIYTYTGKSHFMLPDDDVQ

PGGNLAHAFQPGPGIGGDAHFDEDERWTNNFREYNLHRVAAHELGHSLGLS HST DIGALMYPSYTFSGDVO

LAODD

444

GIQSLYGPGDEDPN GIQAIYGRSQ

FIG. 2A

Sequence 1: MMP-13 Sequence 2: MMP-8

Identity score:

61.4 %

VGEYNVFPRTLKWSKMNLTYRIVNYT PDMTH S EVEKAFKKAFKVWSDVTPLNFTRLHDGIADIMISFGIKEHGDFYPFDG NPKWER T NLTYRIRNYTP QLSEA EVERAI KDAFEL WSVASPLI FTRISQGEADINIAFYQRDHGDNSPFDG

PSGLLAHAFPPGPNYGGDAHFDDDETWTSSSKGYNLFLVAAHEFGHSLGLDHSKDPGALMF <u>PIYTYTGKSHFMLPDDD</u>VQ PNGILAHAFQPGQGIGGDAHFDAEETWTNTSANYNLFLVAA HEFGHSLGLAHSSDPGALMY<u>PNYAF RETSNYSLPODD</u> ID

GIQSLYGPGDEDPN GIQAIYG

FIG. 2B

FIG. 3

| | | Atom Type | Res. | | X | Y . | Z | | |
|------|-----|--------------|------------|--------|---------|------------|---------|------|------|
| MOTA | 1 | | mun | 7 | 10 505 | | | | |
| MOTA | 1 2 | N HN | THR | 7 7 | -12.675 | | -8.815 | 1.00 | 0.83 |
| MOTA | 3 | CA | THR | | -12.001 | | -8.192 | 1.00 | 1.22 |
| MOTA | 4 | HA | THR | · 7 | -14.063 | | -8.340 | 1.00 | 0.63 |
| MOTA | 5 | CB | THR | 7 | | -14.330 | -8.830 | 1.00 | 0.73 |
| MOTA | 6 | HB | THR | 7 | | -13.858 | -6.825 | 1.00 | 0.61 |
| ATOM | 7 | OG1 | THR THR | 7 | | -13.158 | -6.335 | 1.00 | 0.66 |
| ATOM | 8 | HG1 | | 7 | | -15.185 | -6.514 | 1.00 | 0.71 |
| MOTA | 9 | | | 7 | -13.721 | | -7.330 | 1.00 | 1.07 |
| ATOM | 10 | HG21 | | 7 | | -13.628 | -6.336 | 1.00 | 0.67 |
| ATOM | 11 | HG22 | + | 7 | | -12.577 | -6.139 | 1.00 | 1.14 |
| ATOM | 12 | HG23 | THR | 7 | | -14.191 | -5.429 | 1.60 | 1.32 |
| ATOM | 13 | C | | 7 | | -13.955 | -7.093 | 1.00 | 1.23 |
| MOTA | 14 | õ | THR | 7 | | -12.208 | -8.678 | 1.00 | 0.52 |
| MOTA | 15 | N | THR | | | -11.962 | -9.374 | 1.00 | 0.65 |
| MOTA | 16 | | LEU | 8 | | -11.254 | -8.195 | 1.00 | 0.47 |
| MOTA | 17 | HN | LEU | 8 | | -11.473 | -7.639 | 1.00 | 0.61 |
| ATOM | | CA | LEU | 8 | -14.027 | -9.831 | -8.495 | 1.00 | 0.42 |
| | 18 | HA | LEU | 8 | -15.098 | -9.715 | -8.575 | 1.00 | 0.43 |
| MOTA | 19 | | LEU | 8 | -13.495 | -8.937 | -7.370 | 1.00 | 0.52 |
| ATOM | 20 | HB1 | LEU | 8 | -13.721 | -7.905 | -7.591 | 1.00 | 0.54 |
| MOTA | 21 | HB2 | LEU | 8 | -12.424 | -9.060 | -7.292 | 1.00 | 0.58 |
| MOTA | 22 | CG | LEU | 8 | -14.151 | -9.331 | -6.042 | 1.00 | 0.60 |
| MOTA | 23 | HG | LEU | 8 | -13.958 | -10.376 | -5.844 | 1.00 | 0.60 |
| MOTA | 24 | CD1 | | 8 | -13.566 | -8.484 | -4.910 | 1.00 | 0.74 |
| ATOM | 25 | | | 8 | -13.899 | -8.875 | -3.960 | 1.00 | 1.22 |
| ATOM | 26 | | | 8 | -13.900 | -7.462 | -5.016 | 1.00 | 1.26 |
| MOTA | 27 | | | 8 | -12.488 | -8.518 | -4.956 | 1.00 | 1.31 |
| MOTA | 28 | | LEU | 8 | -15.664 | -9.096 | -6.117 | 1.00 | 0.61 |
| MOTA | 29 | | - | 8 | -15.871 | -8.278 | -6.791 | 1.00 | 1.13 |
| ATOM | 30 | | | 8 | -16.040 | -8.856 | -5.134 | 1.00 | 1.18 |
| ATOM | 31 | | | 8 | -16.149 | -9.991 | -6.478 | 1.00 | 1.26 |
| ATOM | 32 | | LEU | 8 8 | -13.374 | -9.438 | -9.822 | 1.00 | 0.40 |
| MOTA | 33 | | LEU | 8 | -12.218 | | -10.064 | 1.00 | 0.45 |
| ATOM | 34 | | LYS | 9 9 | -14.109 | | -10.687 | 1.00 | 0.36 |
| ATOM | 35 | | LYS | 9 | -15.042 | _ | -10.474 | 1.00 | 0.36 |
| ATOM | 36 | | LYS | 9 | | -8.393 | | 1.00 | 0.37 |
| ATOM | 37 | | LYS | 9 | -12.521 | | -11.862 | 1.00 | 0.39 |
| ATOM | 38 | | LYS | 9 | | -9.599 | | 1.00 | 0.50 |
| MOTA | 39 | HB1 | LYS | 9 | -12.851 | -10.344 | -12.573 | 1.00 | 0.60 |

FIG. 4

| | | | | | | . • | | |
|--------|------|--------|--------------|-----------|---------|---------|------|------|
| ATOM | 40 | HB2 LY | S 9 | -13.233 | -9.286 | -13.932 | 1.00 | 0.48 |
| ATOM | 41 | CG LY | | -14.948 | -10.193 | -13.007 | 1.00 | 0.60 |
| ATOM | 42 | HG1 LY | | -15.632 | _ | -13.398 | | |
| | | - | | _ + - + | | | 1.00 | 0.66 |
| MOTA | 43 | HG2 LY | | | | -12.014 | 1.00 | 0.78 |
| ATOM . | . 44 | CD LY | S 9 | · -14.951 | -11.421 | -13.921 | 1.00 | 0.94 |
| ATOM | 45 | HD1 LY | s 9 | -13.944 | -11.794 | -14.033 | 1.00 | 1.57 |
| MOTA | 46 | HD2 LY | | -15.344 | _ | | 1.00 | 1.62 |
| | | | | | | | | |
| ATOM | 47 | CE LY | - | -15.829 | | | 1.00 | 0.57 |
| MOTA | 48 | HE1 LY | S 9 | -16.776 | -12.086 | -13.007 | 1.00 | 1.15 |
| MOTA | 49 | HE2 LY | S 9 | -15.333 | -12.924 | -12.437 | 1.00 | 1.10 |
| ATOM | 50 | NZ LY | | -16.060 | | | 1.00 | 1.61 |
| | | | | | | | | |
| MOTA | 51 | HZ1 LY | | -15.181 | | | 1.00 | 2.14 |
| MOTA | 52 | HZ2 LY | | -16.358 | -13.168 | -15.207 | 1.00 | 2.13 |
| MOTA | 53 | HZ3 LY | s 9 | -16.802 | -14.231 | -13.959 | 1.00 | 2.14 |
| MOTA | 54 | C LY | | -14.377 | | -12.605 | 1.00 | 0.32 |
| MOTA | 55 | O LY | | -15.493 | | -12.191 | | |
| | | | | | | | 1.00 | 0.34 |
| ATOM | 56 | N TR | | -13.850 | | -13.577 | 1.00 | 0.31 |
| MOTA | 57 | HN TR | P 10 | -12.947 | -6.781 | -13.895 | 1.00 | 0.33 |
| ATOM | 58 | CA TR | P 10 | -14.618 | -5.456 | -14.201 | 1.00 | 0.30 |
| ATOM | 59 | HA TR | | -15.030 | - | -13.427 | 1.00 | 0.29 |
| | | | | | | | | |
| MOTA | 60 | CB TR | | -13.684 | | -15.088 | 1.00 | 0.29 |
| ATOM | 61 | HB1 TR | P 10 | -14.264 | -3.917 | -15.655 | 1.00 | 0.32 |
| MOTA | 62 | HB2 TR | P 10 | -13.157 | -5.286 | -15.765 | 1.00 | 0.33 |
| ATOM | 63 | CG TR | | -12.699 | | -14.230 | 1.00 | 0.25 |
| ATOM | 64 | CD1 TR | _ | -11.516 | | | | |
| | | | | | | -13.812 | 1.00 | 0.30 |
| MOTA | 65 | HD1 TR | | -11.137 | -5.390 | -14.040 | 1.00 | 0.37 |
| MOTA | 66 | CD2 TR | P 10 | -12.786 | -2.553 | -13.683 | 1.00 | 0.21 |
| MOTA | 67 | NE1 TR | P 10 | -10.872 | | -13.042 | 1.00 | 0.30 |
| MOTA | 68 | HE1 TR | | -9.996 | | | | |
| | | | | | _ | -12.617 | 1.00 | 0.36 |
| MOTA | 69 | CE2 TF | | -11.614 | | -12.934 | 1.00 | 0.23 |
| MOTA | 70 | CE3 TF | P 10 | -13.758 | -1.538 | -13.763 | 1.00 | 0.24 |
| MOTA | 71 | HE3 TR | P 10 | -14.663 | | -14.328 | 1.00 | 0.29 |
| ATOM | 72 | CZ2 TF | | -11.412 | | -12.287 | | |
| | | _ | | | | | 1.00 | 0.22 |
| MOTA | 73 | HZ2 TF | | -10.509 | | -11.720 | 1.00 | 0.27 |
| MOTA | 74 | CZ3 TF | RP 10 | -13.558 | -0.309 | -13.113 | 1.00 | 0.25 |
| MOTA | 75 | HZ3 TF | RP 10 | -14.310 | 0.463 | -13.181 | 1.00 | 0.32 |
| ATOM | 76 | CH2 TH | | -12.387 | | -12.376 | | |
| | | | | | | | 1.00 | 0.23 |
| MOTA | 77 | HH2 TE | _ · | -12.238 | _ | -11.879 | 1.00 | 0.26 |
| MOTA | 78 | C T | RP 10 | -15.755 | -6.031 | -15.050 | 1.00 | 0.39 |
| MOTA | 79 | O TI | RP 10 | -15.641 | -7.098 | -15.620 | 1.00 | 0.48 |
| MOTA | 80 | N SI | | -16.855 | | -15.132 | 1.00 | 0.43 |
| ATOM | 81 | _ | | | • | | | |
| | | | ER 11 | -16.927 | | -14.660 | 1.00 | 0.44 |
| MOTA | 82 | | ER 11 | -18.006 | -5.835 | -15.936 | 1.00 | 0.52 |
| MOTA | 83 | HA SI | ER 11 | -18.003 | -6.915 | -15.930 | 1.00 | 0.59 |
| MOTA | 84 | CB SI | ER 11 | -19.313 | | -15.325 | 1.00 | 0.64 |
| MOTA | 85 | HB1 SI | | -19.120 | | -14.763 | 1.00 | |
| MOTA | | _ | | | | | | 1.16 |
| | 86 | | ER 11 | -19.718 | | -14.666 | 1.00 | 1.20 |
| ATOM | 87 | OG SI | ER 11 | -20.246 | -5.067 | -16.365 | 1.00 | 1.39 |
| MOTA | 88 | HG SI | ER 11 | -19.821 | -4.495 | -17.008 | 1.00 | 1.92 |
| ATOM | 89 | C SI | ER 11 | -17.893 | | -17.379 | 1.00 | 0.47 |
| ATOM | 90 | _ | ER 11 | | | | | |
| | | | | -18.785 | | -18.181 | 1.00 | 0.60 |
| MOTA | 91 | | rs 12 | -16.808 | | -17.715 | 1.00 | 0.42 |
| MOTA | 92 | HN L | 7S 12 | -16.101 | -4.543 | -17.053 | 1.00 | 0.51 |
| ATOM | 93 | CA L | rs 12 | -16.646 | -4.178 | -19.107 | 1.00 | 0.41 |
| MOTA | 94 | HA L | YS 12 | -17.243 | | -19.781 | 1.00 | 0.47 |
| ATOM | 95 | | rs 12 | -17.116 | | -19.167 | | • |
| ATOM | 96 | | | | | | 1.00 | 0.43 |
| | | , — | | -18.168 | | -18.926 | 1.00 | 0.50 |
| MOTA | 97 | HB2 L | | -16.957 | -2.334 | -20.163 | 1.00 | 0.46 |
| MOTA | 98 | CG L | YS 12 | -16.327 | -1.882 | -18.160 | 1.00 | 0.41 |
| ATOM | 99 | HG1 L | | -15.275 | | -18.401 | 1.00 | 0.37 |
| MOTA | 100 | HG2 L | | -16.484 | | | | |
| ATOM | | | | | . – | -17.164 | 1.00 | 0.42 |
| | 101 | | YS 12 | -16.805 | | -18.223 | 1.00 | 0.50 |
| MOTA | 102 | HD1 L | | -17.856 | -0.386 | -17.981 | 1.00 | 0.56 |
| MOTA | 103 | HD2 L | | -16.648 | | -19.220 | 1.00 | 0.65 |
| ATOM | 104 | | YS 12 | -16.018 | , | -17.218 | | 0.61 |
| ATOM | 105 | HE1 L | | | | | 1.00 | |
| | | | | -15.054 | | -17.636 | | 1.15 |
| ATOM | 106 | HE2 L | | -15.879 | | -16.307 | 1.00 | 1.16 |
| ATOM | 107 | NZ L | YS 12 | -16.773 | 1.661 | -16.920 | 1.00 | 1.39 |
| MOTA | 108 | HZ1 L | | -16.498 | | -15.983 | 1.00 | 1.90 |
| ATOM | 109 | HZ2 L | | | • | | | |
| | | | | -17.794 | | -16.927 | 1.00 | 1.87 |
| ATOM | 110 | | YS 12 | -16.556 | | -17.640 | 1.00 | 1.97 |
| ATOM | 111 | C L | YS 12 | -15.175 | | -19.521 | 1.00 | 0.36 |
| MOTA | 112 | | YS 12 | -14.284 | | -18.695 | | 0.34 |
| ATOM | 113 | _ | ET 13 | -14.917 | | | | |
| ATOM | 114 | | | | | -20.796 | 1.00 | 0.37 |
| | | | ET 13 | -15.652 | | -21.443 | 1.00 | 0.40 |
| ATOM | 115 | | er 13 | -13.506 | _ | -21.269 | 1.00 | 0.38 |
| ATOM | 116 | HA M | ET 13 | -12.910 | | -20.506 | 1.00 | 0.39 |

| ATOM | 117 | CB | MET | 13 | -13.469 | -5.332 -22.543 | 3 1.00 | 0.46 |
|--------|-----|------|-----|------|--------------------|-----------------------|-------------|--------------|
| MOTA | 118 | HB1 | MET | 13 | -12.523 | | | - |
| _ | | | _ | | | -5.189 -23.04 | | 0.53 |
| MOTA | 119 | | | 13 | -14.273 | -5.031 -23.199 | 1.00 | 0.42 |
| MOTA | 120 | CG | MET | 13 | -13.632 | -6.809 - 22.178 | 3 1.00 | 0.64 |
| MOTA | 121 | HG1 | MET | 13 | | | | |
| | | | | | -12.857 | -7.097 - 21.483 | | 1.26 |
| MOTA | 122 | | MET | 13 | -13.556 | -7.411 -23.07 | L 1.00 | 1.37 |
| ATOM | 123 | SD | MET | 13 | -15.252 | -7.067 - 21.414 | 1.00 | 1.22 |
| ATOM | 124 | | MET | 13 | | | | - |
| | | | | | -14.663 | -7.870 -19.90 | | 0.57 |
| MOTA | 125 | HEl | MET | 13 | -14.020 | -7.189 -19.362 | 2 1.00 | 1.16 |
| MOTA | 126 | HE2 | MET | 13 | -14.107 | -8.758 -20.158 | | 1.09 |
| ATOM | 127 | | MET | 13 | | | _ | |
| | | | | | -15.508 | -8.141 - 19.286 | | 1.20 |
| MOTA | 128 | C | MET | 13 | -12.936 | -3.095 -21.560 | 1.00 | 0.32 |
| MOTA | 129 | 0 | MET | 13 | -11.793 | -2.957 - 21.948 | | 0.35 |
| ATOM | 130 | | ASN | 14 | | | | |
| | | | | | -13.718 | -2.064 - 21.37 | | 0.28 |
| MOTA | 131 | HN | ASN | 14 | -14.635 | -2.199 - 21.053 | 2 1.00 | 0.29 |
| MOTA | 132 | CA | ASN | 14 | -13.217 | -0.681 -21.63 | | • |
| ATOM | 133 | | ASN | | | | | 0.26 |
| | | | | 14 | -12.359 | -0.725 -22.286 | | 0.29 |
| MOTA | 134 | | ASN | 14 | -14.319 | 0.148 -22.29 | 7 1.00 | 0.30 |
| MOTA | 135 | HB1 | ASN | 14 | -14.025 | 1.186 -22.318 | | 0.31 |
| MOTA | 136 | | ASN | | | | | |
| | | | | 14 | -15.235 | 0.043 -21.73 | | 0.31 |
| MOTA | 137 | | asn | 14 | -14.539 | -0.346 - 23.729 | 9 1.00 | 0.37 |
| MOTA | 138 | OD1 | ASN | 14 | -13.677 | -0.981 -24.304 | | 1.16 |
| ATOM | 139 | | ASN | 14 | | | | |
| | | | | | -15.664 | -0.077 -24.334 | | 1.05 |
| MOTA | 140 | | | 14 | -16.359 | 0.435 -23.87 | 1.00 | 1.81 |
| MOTA | 141 | HD22 | asn | 14 | -15.812 | -0.386 -25.25 | | 1.06 |
| MOTA | 142 | C | ASN | 14 | -12.813 | | | |
| ATOM | 143 | | | | | | | 0.22 |
| | | | ASN | 14 | -13.566 | -0.019 -19.35 | 7 1.00 | 0.23 |
| MOTA | 144 | N | LEU | 15 | -11.630 | 0.533 -20.24 | | 0.21 |
| MOTA | 145 | HN | LEU | 15 | -11.042 | | | |
| ATOM | 146 | | | | | 0.517 -21.03 | | 0.24 |
| | | | LEU | 15 | -11.171 | 1.194 -18.98 | 7 1.00 | 0.18 |
| MOTA | 147 | HA | LEU | 15 | -12.025 | 1.447 -18.37 | 9 1.00 | 0.19 |
| ATOM | 148 | CB | LEU | 15 | -10.250 | 0.243 -18.210 | | |
| MOTA | 149 | | | | • - • | | | 0.18 |
| | | | LEU | 15 | -9.812 | 0.769 -17.37 | 5 1.00 | 0.19 |
| ATOM | 150 | HB2 | LEU | 15 | -9.463 | -0.102 -18.86 | 5 1.00 | 0.21 |
| MOTA | 151 | CG | LEU | 15 | -11.046 | -0.964 -17.69 | | |
| ATOM | 152 | | LEU | | | | | 0.19 |
| | | _ | | 15 | -11.547 | -1.442 - 18.52 | + | 0.20 |
| ATOM | 153 | CD1 | LEU | 15 | -10.086 | -1.961 -17.04 | 1.00 | 0.20 |
| ATOM | 154 | HD11 | LEU | 15 | -9.726 | -1.556 -16.11 | | |
| MOTA | 155 | | | | | | | 0.98 |
| | | | | 15 | -9.251 | -2.141 -17.70 | | 1.04 |
| MOTA | 156 | | LEU | 15 | -10.604 | -2.890 -16.85 | 7 1.00 | 1.07 |
| ATOM | 157 | CD2 | LEU | 15 | -12.083 | -0.513 -16.65 | | |
| MOTA | 158 | | | 15 | | | | 0.21 |
| | | | _ | | -12.114 | -1.228 - 15.85 | | 1.07 |
| MOTA | 159 | | LEU | 15 | -13.055 | -0.456 - 17.12 | 2 1.00 | 1.00 |
| MOTA | 160 | HD23 | LEU | 15 | -11.814 | 0.457 -16.26 | | 1.04 |
| MOTA | 161 | C | LEU | 15 | | | | |
| ATOM | | _ | | | -10.397 | 2.471 - 19.33 | | 0.18 |
| | 162 | | LEU | - 15 | -9.785 | 2.570 -20.38 | 1.00 | 0.20 |
| ATOM . | 163 | N | THR | 16 | -10.425 | 3.447 -18.46 | | 0.18 |
| MOTA | 164 | HN | THR | 16 | -10.929 | 3.338 -17.62 | _ : : : : : | |
| MOTA | 165 | | THR | 16 | | | • • | 0.18 |
| | | | | | -9.699 | 4.729 -18.72 | 2 1.00 | 0.19 |
| MOTA | 166 | | THR | 16 | -9.051 | 4.617 -19.57 | 1.00 | 0.20 |
| MOTA | 167 | CB | THR | 16 | -10.716 | 5.839 -18.99 | | |
| MOTA | 168 | HB | THR | 16 | | | | 0.22 |
| ATOM | | | | | -10.198 | 6.729 - 19.31 | | 0.24 |
| | 169 | | | 16 | -11.445 | 6.112 -17.80 | 3 1.00 | 0.23 |
| MOTA | 170 | HG1 | THR | 16 | -11.821 | · 5.286 -17.49 | 1.00 | 0.98 |
| MOTA | 171 | CG2 | THR | 16 | -11.680 | 5.393 -20.09 | | |
| MOTA | | HG21 | | | | | | 0.26 |
| | | | | 16 | -12.200 | 6.254 - 20.48 | | 1.05 |
| ATOM | | HG22 | THR | 16 | -12.396 | 4.696 -19.686 | 5 1.00 | 1.02 |
| ATOM | 174 | HG23 | THR | 16 | -11.125 | 4.914 -20.889 | | 1.05 |
| MOTA | 175 | | THR | 16 | -8.864 | 5.100 -17.49 | | |
| ATOM | 176 | - | | | | | _ • • • | 0.17 |
| | | _ | THR | 16 | - 9.157 | 4.687 -16.39 | 1.00 | 0.16 |
| MOTA | 177 | N | TYR | 17 | -7.826 | 5.878 -17.679 | | 0.18 |
| ATOM | 178 | HN | TYR | 17 | -7.603 | 6.202 -18.57 | | |
| MOTA | 179 | | | | | | • • • | 0.19 |
| | | | TYR | 17 | -6.981 | 6.268 - 16.50 | | 0.17 |
| ATOM . | 180 | | TYR | 17 | - 7.585 | 6.233 -15.61 | 5 1.00 | 0.17 |
| MOTA | 181 | CB | TYR | 17 | -5.814 | 5.288 -16.36 | | |
| ATOM | 182 | | | 17 | | | | 0.19 |
| MOTA | | | | | -6.194 | 4.278 -16.34 | | 0.19 |
| | 183 | | TYR | 17 | -5.292 | 5.488 -15.438 | | 0.20 |
| MOTA | 184 | | TYR | 17 | -4.857 | 5.445 -17.520 | | 0.22 |
| ATOM | 185 | | | 17 | | | | |
| ATOM | 186 | | | | -5.037 | 4.685 -18.682 | | 0.26 |
| | | | TYR | 17 | -5.867 | 3.998 -18.75 | 1.00 | 0.27 |
| ATOM | 187 | | TYR | 17 | -3.782 | 6.336 -17.426 | | 0.25 |
| ATOM | 188 | HD2 | TYR | 17 | -3.643 | | | |
| ATOM | 189 | | | 17 | | 6.923 -16.530 | | 0.26 |
| | | | | | -4.143 | 4.817 -19.75 | | 0.31 |
| ATOM | 190 | | | 17 | -4.282 | 4.231 -20.643 | | 0.36 |
| MOTA | 191 | CE2 | TYR | 17 | -2.888 | 6.470 -18.49 | | 0.30 |
| ATOM | 192 | HE2 | TYR | 17 | -2.059 | | | |
| ATOM | 193 | | TYR | | | 7.158 -18.424 | | 0.35 |
| | 133 | CZ | TIK | 17 | -3.068 | 5.710 - 19.658 | 3 1.00 | 0.32 |
| | | | | | | | | |

| a mov | 104 Au mun | • • | . 0 100 | | |
|--------------|--------------------------|------|--------------------|--------------------------------|------------------------|
| MOTA | 194 OH TYR 195 HH TYR | 17 | -2.186 | 5.839 -20.711 | 1.00 0.39 |
| MOTA | | 17 | -1.696 | 5.016 -20.790 | 1.00 0.85 |
| MOTA | | 17 | -6.448 | 7.692 -16.690 | 1.00 0.19 |
| MOTA | 197 O TYR | 17 | -6.414 | 8.220 -17.784 | 1.00 0.21 |
| MOTA | 198 N ARG | 18 | -6.044 | 8.320 -15.616 | 1.00 0.19 |
| MOTA | 199 HN ARG | 18 | -6.089 | 7.874 -14.747 | 1.00 0.19 |
| MOTA | 200 CA ARG | 18 | -5.523 | 9.714 - 15.712 | 1.00 0.22 |
| MOTA | 201 HA ARG | 18 | -5.131 | 9.877 -16.704 | 1.00 0.24 |
| ATOM | 202 CB ARG | 18 | -6.674 | 10.691 -15.447 | 1.00 0.27 |
| MOTA | 203 HB1 ARG | 18 | -6.978 | 10.613 -14.412 | 1.00 0.31 |
| MOTA | 204 HB2 ARG | 18 | - 7.507 | 10.442 -16.083 | 1.00 0.30 |
| ATOM | 205 CG ARG | 18 | -6.229 | 12.127 -15.733 | 1.00 0.35 |
| MOTA | 206 HG1 ARG | 18 | -5.504 | 12.137 -16.531 | 1.00 0.93 |
| MOTA | 207 HG2 ARG | 18 | -5.790 | 12.549 -14.843 | 1.00 0.85 |
| MOTA | 208 CD ARG | 18 | -7.447 | 12.946 -16.149 | 1.00 0.81 |
| ATOM | 209 HD1 ARG | 18 | -8.216 | 12.867 -15.378 | 1.00 1.29 |
| MOTA | 210 HD2 ARG | 18 | -7.838 | 12.561 ~17.068 | 1.00 1.63 |
| MOTA | 211 NE ARG | 18 | -7.030 | 14.362 -16.406 | 1.00 1.52 |
| MOTA | 212 HE ARG | 18 | -7.071 | 14.711 -17.318 | 1.00 2.11 |
| MOTA | 213 CZ ARG | 18 | -6.561 | 15.119 -15.456 | 1.00 2,24 |
| MOTA | 214 NH1 ARG | · 18 | -6.119 | 16.314 -15.736 | 1.00 3.18 |
| MOTA | 215 HH11 ARG | 18 | -6.142 | 16.647 -16.679 | 1.00 3.48 |
| MOTA | 216 HH12 ARG | 18 | -5.760 | 16.898 -15.009 | 1.00 3.84 |
| MOTA | 217 NH2 ARG | 18 | -6.564 | 14.700 -14.220 | 1.00 2.63 |
| ATOM . | 218 HH21 ARG | 18 | -6.928 | 13.795 -14.000 | 1.00 2.44 |
| MOTA | 219 HH22 ARG | 18 | -6.205 | 15.285 -13.493 | 1.00 3.49 |
| MOTA | 220 C ARG | 18 | -4.413 | 9.931 -14.676 | 1.00 0.21 |
| MOTA | 221 O ARG | 18 | -4.550 | 9.576 -13.522 | 1.00 0.23 |
| MOTA | 222 N ILE | 19 | -3.314 | 10.514 -15.079 | 1.00 0.21 |
| MOTA | 223 HN ILE | 19 | -3.223 | 10.794 -16.014 | 1.00 0.22 |
| ATOM | 224 CA ILE | 19 | -2.196 | 10.755 -14.118 | 1.00 0.23 |
| MOTA | 225 HA ILE | 19 | -2.200 | 9.985 -13.360 | 1.00 0.25 |
| MOTA | 226 CB ILE | | -0.864 | 10.721 -14:875 | 1.00 0.25 |
| ATOM | 227 HB ILE | | -0.862 | 11.491 -15.633 | 1.00 0.25 |
| ATOM | 228 CG1 ILE | | -0.702 | 9.341 -15.531 | 1.00 0.29 |
| ATOM | 229 HG11 ILE | | -1.607 | 9.092 -16.065 | 1.00 0.82 |
| MOTA | 230 HG12 ILE | | -0.525 | 8.601 -14.765 | 1.00 0.97 |
| ATOM | 231 CG2 ILE | | 0.291 | 10.962 -13.893 | 1.00 0.29 |
| MOTA | 232 HG21 ILE | -• | 1.231 | 10.914 -14.420 | 1.00 1.08 |
| ATOM | 233 HG22 ILE | | 0.272 | 10.206 -13.123 | 1.00 1.09 |
| ATOM | 234 HG23 ILE | - • | 0.187 | 11.937 -13.440 | 1.00 1.00 |
| ATOM | 235 CD1 ILE | | 0.477 | 9.345 -16.512 | 1.00 0.93 |
| ATOM | 236 HD11 ILE | | 1.402 | 9.216 -15.970 | 1.00 1.59 |
| MOTA | 237 HD12 ILE | | 0.501 | 10.280 -17.050 | 1.00 1.50 |
| ATOM | 238 HD13 ILE | _ | 0.360 | 8.533 -17.214 | 1.00 1.55 |
| MOTA MOTA | 239 C ILE | _ | -2.381 | 12.126 -13.454 | 1.00 0.23 |
| ATOM | 240 O ILE | _ | -2.355 | 13.150 -14.108 | 1.00 0.23 |
| ATOM | 241 N VAL 242 HN VAL | | -2.563 | 12.152 -12.161 | 1.00 0.25 |
| ATOM | | - • | -2.578 | 11.314 -11.653 | 1.00 0.27 |
| MOTA | 243 CA VAL 244 HA VAL | | -2.746 | 13.454 -11.454 | 1.00 0.27 |
| ATOM | 245 CB VAL | | -3.496 | 14.035 -11.970 | 1.00 0.27 |
| ATOM | 246 HB VAL | | - 3.202 | 13.205 -10.015 | 1.00 0.31 |
| ATOM | 247 CG1 VAL | | -2.522 -3.216 | 12.517 -9.534 | 1.00 0.32 |
| ATOM | 248 HG11 VAL | _ | -3.607 | 14.529 -9.247 | 1.00 0.33 |
| ATOM | 249 HG12 VAL | | -2.211 | 15.310 -9.883 14.782 -8.944 | 1.00 0.97 |
| MOTA | 250 HG13 VAL | _ | -3.842 | 14.782 -8.944 14.432 -8.372 | 1.00 1.08 |
| MOTA | 251 CG2 VAL | | -4.612 | 12.611 -10.028 | 1.00 1.10 1.00 0.33 |
| ATOM | 252 HG21 VAL | | -5.296 | 13.317 -10.476 | 1.00 1.05 |
| ATOM | 253 HG22 VAL | | -4.924 | 12.401 -9.016 | 1.00 1.03 |
| MOTA | 254 HG23 VAL | | -4.612 | 11.697 -10.602 | 1.00 1.03 |
| MOTA | 255 C VAL | | -1.424 | 14.231 -11.451 | 1.00 0.27 |
| MOTA | 256 O VAL | | -1.403 | 15.435 -11.611 | 1.00 0.26 |
| MOTA | 257 N ASN | | -0.321 | 13.555 -11.259 | 1.00 0.28 |
| MOTA | 258 HN ASN | | -0.357 | 12.585 -11.124 | 1.00 0.30 |
| ATOM | 259 CA ASN | | 0.992 | 14.265 -11.235 | 1.00 0.29 |
| MOTA | 260 HA ASN | | 0.973 | 15.076 -11.949 | 1.00 0.26 |
| MOTA | 261 CB ASN | | 1.235 | 14.829 -9.834 | 1.00 0.33 |
| MOTA | 262 HB1 ASN | | 0.544 | 15.637 -9.646 | 1.00 0.33 |
| ATOM | 263 HB2 ASN | | 2.249 | 15.199 -9.766 | 1.00 0.35 |
| MOTA | 264 CG ASN | | 1.022 | 13.727 -8.795 | 1.00 0.40 |
| ATOM | 265 OD1 ASN | 21 | 0.459 | 12.694 -9.097 | 1.00 1.01 |
| MOTA | 266 ND2 ASN | 21 | 1.445 | 13.908 -7.574 | 1.00 0.88 |
| MOTA | 267 HD21 ASN | | 1.895 | 14.743 -7.330 | 1.00 1.50 |
| MOTA | 268 HD22 ASN | 21 | 1.312 | 13.208 -6.901 | 1.00 0.88 |
| ATOM | 269 C ASN | | 2.116 | 13.291 -11.606 | 1.00 0.34 |
| MOTA | 270 o asn | 21 | 1.929 | 12.090 -11.619 | 1.00 0.37 |
| | | | | | |

| MOTA | 271 | N | TYR | 22 | 3.274 | 13.810 -11. | 933 1.00 | 0.38 |
|-------|-----|------|-------|------|--------|-------------|-----------|--------------|
| ATOM | 272 | | TYR | 22 | | | | |
| | | | | | 3.387 | 14.783 -11. | | 0.38 |
| ATOM | 273 | | TYR | 22 | 4.417 | 12.935 -12. | | 0.46 |
| ATOM | 274 | HA | TYR | 22 | 4.067 | 11.929 -12. | 509 1.00 | 0.45 |
| ATOM | 275 | CB | TYR | 22 | 5.028 | 13.481 -13. | 630 1.00 | 0.49 |
| ATOM | 276 | HB1 | TYR | 22 | 5.845 | 12.846 -13. | | |
| ATOM | 277 | | TYR | 22 | 5.397 | 14.482 -13. | | |
| | | | | | | | | |
| ATOM | 278 | | TYR | 22 | 3.981 | 13.513 -14. | | 0.43 |
| MOTA | 279 | | TYR | 22 | 3.684 | 12.352 -15. | 436 1.00 | 0.38 |
| MOTA | 280 | HD1 | TYR | 22 | 4.199 | 11.430 -15. | 212 1.00 | 0.39 |
| ATOM | 281 | CD2 | TYR | 22 | 3.313 | 14.708 -15. | | |
| ATOM | 282 | | TYR | 22 | 3.543 | 15.603 -14. | | |
| ATOM | 283 | | TYR | 22 | 2.718 | 12.386 -16. | | |
| MOTA | | | | | | | | - |
| | 284 | | TYR | 22 | 2.490 | 11.491 -17. | | |
| MOTA | 285 | | TYR | 22 | 2.345 | 14.742 -16. | 013 1.00 | 0.44 |
| MOTA | 286 | HE2 | TYR | 22 | 1.828 | 15.663 -16. | 235 1.00 | 0.49 |
| ATOM | 287 | CZ | TYR | 22 | 2.048 | 13.581 -16. | 735 1.00 | |
| ATOM | 288 | OH | TYR | 22 | 1.095 | 13.615 -17. | | _ |
| MOTA | 289 | | TYR | 22 | 1.173 | 14.457 -18. | | |
| ATOM | 290 | | TYR | 22 | | | | |
| | | | | | 5.499 | 12.923 -11. | | |
| MOTA | 291 | | TYR | 22 | 6.554 | 12.378 -11. | 470 1.00 | 1.38 |
| MOTA | 292 | N | THR | 23 | 5.240 | 13.544 -10. | 130 1.00 | 0.47 |
| ATOM | 293 | HN | THR | 23 | 4.372 | 13,987 -10. | 023 1.00 | |
| ATOM | 294 | CA | THR | 23 | 6.237 | | 004 1.00 | |
| MOTA | 295 | HA | THR | 23 | 5.848 | | | |
| ATOM | 296 | | | | | | 304 1.00 | |
| | | | THR | 23 | 6.361 | | 273 1.00 | |
| MOTA | 297 | HB | THR . | - • | 5.383 | 11.969 -7. | .921 1.00 | 0.68 |
| MOTA | 298 | OG1 | THR | 23 | 7.223 | 12.420 -7. | 156. 1.00 | 0.86 |
| ATOM | 299 | HG1 | THR | 23 | 7.941 | | 244 1.00 | |
| MOTA | 300 | CG2 | THR | 23 | 6.916 | | | |
| ATOM | 301 | | | 23 | | | | |
| | | | | | 7.753 | | .748 1.00 | |
| MOTA | | HG22 | | 23 | 6.141 | | .850 1.00 | 1.16 |
| ATOM | 303 | HG23 | THR | 23 | 7.245 | 10.332 - 8 | .570 1.00 | 1.22 |
| MOTA | 304 | C | THR | 23 | 7.623 | 14.115 -9 | .523 1.00 | |
| MOTA | 305 | 0 | THR | 23 | 8.077 | 13.699 -10. | | |
| MOTA | 306 | N | PRO | 24 | 8.302 | | | • |
| MOTA | 307 | | | | | | 823 1.00 | |
| | | CA | PRO | 24 | 9.625 | | .311 1.00 | - |
| MOTA | 308 | HA | PRO | 24 | 9.534 | | .307 1.00 | 0.46 |
| ATOM | 309 | CB | PRO | 24 | 9.924 | 16.655 -8 | .335 1.00 | 0.50 |
| ATOM | 310 | HB1 | PRO | 24 | 9.743 | | .815 1.00 | |
| MOTA | 311 | HB2 | PRO | 24 | 10.955 | | .014 1.00 | |
| ATOM | 312 | CG | PRO | 24 | 8.995 | | | – – |
| ATOM | 313 | | | | | • | .129 1.00 | |
| | | HG1 | PRO | 24 | 8.613 | | .842 1.00 | |
| MOTA | 314 | HG2 | PRO | 24 | 9.537 | 16.069 - 6 | .303 1.00 | 0.76 |
| MOTA | 315 | CD | PRO | 24 | 7.832 | 15.598 -7 | .529 1.00 | 0.56 |
| MOTA | 316 | HD2 | PRO | 24 | 7.675 | | .786 1.00 | |
| MOTA | 317 | HD1 | PRO | 24 | 6.940 | | .680 1.00 | |
| MOTA | 318 | C | PRO | | | | | |
| | | | | 24 | 10.743 | | .253 1.00 | |
| ATOM | 319 | 0 | PRO | 24 | 11.835 | | .737 1.00 | 0.40 |
| MOTA | 320 | N | ASP | 25 | 10.490 | 13.337 -8 | .662 1.00 | 0.44 |
| ATOM | 321 | HN | ASP | 25 | 9.608 | 13.172 -8 | .270 1.00 | |
| ATOM | 322 | CA | ASP | 25 | 11.554 | | .577 1.00 | |
| ATOM | 323 | HA | ASP | 25 | 12.393 | | | |
| ATOM | 324 | | | | | | .025 1.00 | |
| _ | | CB | ASP | 25 | 11.016 | | .847 1.00 | |
| ATOM | 325 | | ASP | 25 | 11.719 | | .945 1.00 | |
| ATOM | 325 | | ASP | 25 | 10.068 | 10.773 -8 | .276 1.00 | 0.56 |
| MOTA | 327 | CG | ASP | 25 | 10.827 | | .364 1.00 | |
| ATOM | 328 | OD1 | ASP | 25 | 10.079 | | .709 1.00 | |
| ATOM | 329 | | ASP | 25 | 11.437 | | | |
| ATOM | 330 | C | | | | | | |
| | | | ASP | 25 | 12.025 | | .985 1.00 | |
| MOTA | 331 | 0 | ASP | 25 | 13.179 | 11.597 -10 | | 0.55 |
| MOTA | 332 | N | MET | 26 | 11.146 | 11.948 -10 | .955 1.00 | |
| MOTA | 333 | HN | MET | 26 | 10.220 | 12.209 -10 | | |
| ATOM | 334 | CA | MET | 26 | 11.553 | 11.590 -12 | | |
| ATOM | 335 | HA | MET | 26 | | | | |
| MOTA | | | | | 12.624 | 11.686 -12 | | |
| | 336 | | MET | 26 | 11.144 | 10.149 -12 | | |
| MOTA | 337 | | MET | 26 | 11.282 | 9.954 -13 | .709 1.00 | 0.55 |
| MOTA | 338 | HB2 | MET | . 26 | 10.105 | 10.006 -12 | | |
| ATOM | 339 | CG | MET | 26 | 12.011 | 9.186 -11 | | |
| ATOM | 340 | | MET | 26 · | 11.783 | 9.288 -10 | | _ |
| ATOM | 341 | | | | | | | |
| | | | MET | 26 | 13.053 | 9.419 -12 | . | |
| MOTA | 342 | SD | MET | 26 | 11.683 | 7.485 - 12 | | 0.89 |
| ATOM | 343 | CE. | | 26 | 10.000 | 7.330 -11 | | |
| ATOM | 344 | HE1 | MET | 26 | 9.292 | 7.456 -12 | | |
| MOTA | 345 | | MET | 26 | 9.825 | 8.084 -10 | | · — - |
| ATOM | 346 | HE3 | MET | 26 | 9.877 | | | |
| ATOM | _ | | | | | 6.352 -11 | | |
| WA OU | 347 | С | MET | 26 | 10.872 | 12.530 -13 | .344 1.00 | 0.34 |

| ATOM | 348 | 0 | MET | 26 | 9.897 | 13.184 -13.031 | 1.00 | 0.32 |
|--|---|---|--|--|--|---|--|--|
| ATOM | 349 | N | THR | 27 | 11.385 | 12.604 -14.544 | | |
| ATOM | 350 | | | | | | 1.00 | 0.33 |
| | | HN | THR | 27 | 12.174 | 12.070 -14.773 | 1.00 | 0.38 |
| ATOM | 351 | CA | THR | 27 | 10.775 | 13.504 -15.562 | 1.00 | 0.32 |
| MOTA | 352 | HA | THR | 27 | 10.618 | 14.483 -15.133 | 1.00 | 0.35 |
| MOTA | 353 | CB | THR | 27 | 11.711 | 13.616 -16.768 | 1.00 | 0.39 |
| ATOM | 354 | HB | THR | 27 | | | | |
| | | | | | 11.295 | 14.308 -17.484 | 1.00 | 0.42 |
| MOTA | 355 | OG1 | THR | 27 | 11.852 | 12.338 -17.371 | 1.00 | 0.37 |
| MOTA | 356 | HG1 | THR | .27 | 12.765 | 12.242 -17.653 | 1.00 | 0.94 |
| MOTA | 357 | CG2 | THR | 27 | 13.080 | 14.121 -16.313 | 1.00 | 0.51 |
| MOTA | 358 | HG21 | THR | 27 | 13.602 | _ | | |
| MOTA | 359 | | | | | 14.553 -17.154 | 1.00 | 1.14 |
| | | HG22 | THR | 27 | 13.655 | 13.297 -15.918 | 1.00 | 1.11 |
| MOTA | 360 | HG23 | THR | 27 | 12.951 | 14.871 -15.546 | 1.00 | 1.12 |
| MOTA | 361 | C . | THR | 27 | 9.436 | 12.921 -16.013 | 1.00 | 0.27 |
| ATOM | 362 | 0 | THR | 27 | 9.177 | 11.743 -15.864 | 1.00 | 0.24 |
| MOTA | 363 | N | HIS | 28 | 8.580 | | | |
| ATOM | 364 | | | | | 13.740 -16.554 | 1.00 | 0.32 |
| | | HN | HIS | 28 | 8.807 | 14.688 -16.657 | 1.00 | 0.37 |
| MOTA | 365 | · CA | HIS | 28 | 7.253 | 13.241 -17.004 | 1.00 | 0.34 |
| MOTA | 366 | HA | HIS | 28 | 6.715 | 12.833 -16.161 | 1.00 | 0.36 |
| MOTA | 367 | CB | HIS | 28 | 6.457 | 14.403 -17.601 | 1.00 | 0.46 |
| MOTA | 368 | HB1 | HIS | 28 | 5.428 | _ | | |
| ATOM | 369 | | | | | 14.104 -17.736 | 1.00 | 0.71 |
| | | HB2 | HIS | 28 | 6.880 | 14.676 -18.557 | 1.00 | 0.88 |
| ATOM | 370 | CG | HIS | 28 | 6.516 | 15.583 -16.669 | 1.00 | 0.73 |
| ATOM | 371 | ND1 | HIS | 28 . | 6.056 | 16.838 -17.036 | 1.00 | 1.66 |
| MOTA | 372 | HD1 | HIS | 28 | 5.659 | 17.080 -17.898 | 1.00 | 2.30 |
| MOTA | 373 | CD2 | | 28 | 6.987 | 15.716 -15.387 | 1.00 | |
| ATOM | 374 | | HIS | 28 | | | | 1.33 |
| ATOM | | | | | 7.423 | 14.922 -14.798 | 1.00 | 2.01 |
| | 375 | | HIS | 28 | 6.258 | 17.664 -15.993 | 1.00 | 1.95 |
| ATOM | 376 | HEl | HIS | 28 | 5.993 | 18.711 -15.990 | 1.00 | 2.70 |
| MOTA | 377 | NE2 | HIS | 28 | 6.823 | 17.031 -14.962 | 1.00 | 1.71 |
| ATOM | 378 | C | HIS | 28 | 7.436 | 12.156 -18.069 | | _ |
| MOTA | 379 | ŏ | | | | | 1.00 | 0.30 |
| | | | HIS | 28 | 6.737 | 11.164 -18.082 | 1.00 | 0.30 |
| ATOM | 380 | N | SER | 29 | 8.362 | 12.338 -18.970 | 1.00 | 0.31 |
| ATOM | 381 | HN | SER | 29 | 8.912 | 13.149 -18.952 | 1.00 | 0.34 |
| ATOM | 382 | CA | SER | 29 | 8.567 | 11.319 -20.039 | 1.00 | 0.32 |
| MOTA | 383 | HA | SER | 29 | 7.660 | 11.217 -20.615 | | |
| ATOM | 384 | СВ | SER | | | | 1.00 | 0.35 |
| | | | _ | . 29 | 9.699 | 11.775 -20.959 | 1.00 | 0.38 |
| MOTA | 385 | _ | SER | 29 | 9.973 | 10.963 -21.621 | 1.00 | 0.39 |
| ATOM | 386 | HB2 | SER | 29 | 10.555 | 12.056 -20.368 | 1.00 | 0.37 |
| MOTA | 387 | OG | SER | 29 | 9.265 | 12.896 -21.717 | 1.00 | 0.45 |
| MOTA | 388 | HG | SER | 29 | 9.157 | 12.614 -22.628 | 1.00 | 0.96 |
| ATOM | 389 | | SER | 29 | 8.931 | 9.964 -19.424 | | |
| MOTA | 390 | | SER | 29 | | | 1.00 | 0.26 |
| MOTA | 391 | - | | | 8.479 | 8.930 -19.876 | 1.00 | 0.26 |
| | | N | GLU | 30 | 9.747 | 9.954 -18.405 | 1.00 | 0.24 |
| ATOM | 392 | HN | GLU | 30 | 10.107 | 10.796 -18.056 | 1.00 | 0.25 |
| ATOM | 393 | CA | GLU | 30 | 10.137 | 8.657 -17.779 | 1.00 | 0.22 |
| MOTA | 394 | HA | GLU | 30 | 10.484 | 7.978 -18.542 | 1.00 | 0.25 |
| MOTA | 395 | CB | GLU | 30 | 11.260 | | | |
| MOTA | 396 | | GLU | | | 8.899 -16.769 | | 0.23 |
| | | | | 30 | 11.424 | 8.002 -16.191 | 1.00 | 0.24 |
| ATOM | 397 | HB2 | GLU | 30 | 10.980 | 9.707 -16.108 | 1.00 | 0.22 |
| ATOM | 398 | CG | GLŲ | 30 | 12.547 | 9.268 -17.510 | 1.00 | 0.29 |
| MOTA | 399 | HG1 | GLŲ | 30 | 12.386 | 10.165 -18.086 | 1.00 | 0.67 |
| ATOM | 400 | HG2 | GLU | 30 | 12.826 | 8.460 -18.171 | 1.00 | 0.68 |
| MOTA | 401 | CD | GLU | 30 | 13.666 | 9.509 -16.495 | | |
| ATOM | 402 | | | | | | 1.00 | 0.84 |
| MOTA | | | GLU | 30 | 13.436 | 9.266 -15.321 | 1.00 | 1.49 |
| | 403 | | | 30 | 14.731 | 9:936 -16.908 | 1.00 | 1.59 |
| MOTA | 404 | C | GLU | 30 | 8.935 | 8.046 -17.051 | 1.00 | 0.17 |
| MOTA | 405 | | GLU | 30 | 8.715 | 6.849 -17.082 | 1.00 | 0.19 |
| MOTA | 406 | N | VAL | 31 | 8.163 | 8.861 -16.387 | 1.00 | |
| MOTA | 407 | | VAL | 31 | 8.366 | | | 0.16 |
| MOTA | 408 | | | | | 9,819 -16.371 | 1.00 | 0.17 |
| | | | VAL | 31 | 6.983 | 8.341 -15.640 | 1.00 | 0.16 |
| ATOM | 409 | | VAL | 31 | 7.292 | 7.527 -14.999 | 1.00 | 0.17 |
| MOTA | | ~~ | VAL | 31 | 6.402 | 9.464 -14.782 | | |
| ATOM | 410 | | AVD | | | 21102 721112 | 1.00 | U.20 |
| MOTA | 410 | | | | | | | 0.20 |
| | 411 | HB | VAL | 31 | 6.261 | 10,344 -15.392 | 1.00 | 0.22 |
| | 411 412 | HB CG1 | VAL VAL | 31 31 | 6.261 5.058 | 10,344 -15.392 9.021 -14.208 | 1.00 | 0.22 0.23 |
| MOTA | 411 412 413 | HB CG1 HG11 | VAL VAL | 31 31 31 | 6.261 5.058 5.135 | 10.344 -15.392 9.021 -14.208 8.000 -13.867 | 1.00 1.00 1.00 | 0.22 0.23 0.97 |
| MOTA MOTA | 411 412 413 414 | HB CG1 HG11 HG12 | VAL VAL VAL | 31 31 31 31 | 6.261 5.058 5.135 4.298 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 | 1.00 | 0.22 0.23 |
| MOTA MOTA MOTA | 411 412 413 414 415 | HB CG1 HG11 HG12 HG13 | VAL VAL VAL VAL | 31 31 31 31 31 | 6.261 5.058 5.135 | 10.344 -15.392 9.021 -14.208 8.000 -13.867 | 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 |
| MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 | HB CG1 HG11 HG12 HG13 CG2 | VAL VAL VAL | 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 | 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 |
| MOTA MOTA MOTA | 411 412 413 414 415 | HB CG1 HG11 HG12 HG13 CG2 | VAL VAL VAL VAL VAL | 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 | 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 |
| MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 | HB CG1 HG11 HG12 HG13 CG2 HG21 | VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 | 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 |
| MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 | VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 | 10.344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 | 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 |
| MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 | VAL VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 | 1.00 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 |
| MOTA MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 C | VAL VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 5.911 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9:090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 7.844 -16.617 | 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 |
| MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 420 421 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 C | VAL | 31 31 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 | 1.00 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 0.99 0.16 |
| MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 420 421 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 C | VAL VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 5.911 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 7.844 -16.617 6.817 -16.406 | 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 0.99 0.16 0.17 |
| MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 420 421 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 C | VAL | 31 31 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 5.911 5.293 5.672 | 10.344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 7.844 -16.617 6.817 -16.406 8.571 -17.677 | 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 0.99 0.16 0.17 |
| MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA | 411 412 413 414 415 416 417 418 419 420 421 | HB CG1 HG11 HG12 HG13 CG2 HG21 HG22 HG23 C O N HN | VAL VAL VAL VAL VAL VAL VAL VAL | 31 31 31 31 31 31 31 31 31 | 6.261 5.058 5.135 4.298 4.793 7.364 7.528 6.936 8.304 5.911 5.293 | 10,344 -15.392 9.021 -14.208 8.000 -13.867 9.090 -14.973 9.659 -13.378 9.785 -13.636 8.897 -13.045 10.557 -13.013 10.129 -14.040 7.844 -16.617 6.817 -16.406 | 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 | 0.22 0.23 0.97 1.07 1.07 0.24 1.05 1.03 0.99 0.16 0.17 |

| MOTA | 425 | HA GLU | 32 | . 3.673 | 9 002 -19 147 | 1 00 | 0 04 |
|------|-------|---------|----|---------|----------------|------|------|
| MOTA | 426 | | | • | 8.092 -18.147 | | 0.24 |
| | | CB GLU | 32 | 4.533 | 9.170 -19.787 | 1.00 | 0.27 |
| MOTA | 427 | HB1 GLU | 32 | 3.922 | 8.772 -20.582 | 1.00 | 0.31 |
| MOTA | 428 | HB2 GLU | 32 | 5.524 | 9.379 -20.164 | 1.00 | 0.28 |
| MOTA | 429 | CG GLÜ | 32 | 3.904 | 10.463 -19.262 | | 0.29 |
| MOTA | 430 | HG1 GLU | 32 | 4.456 | | | |
| MOTA | 431 | _ | | | 10.812 -18.405 | | 0.48 |
| | | | 32 | 2.879 | 10.272 -18.977 | | 0.52 |
| MOTA | 432 | CD GLU | 32 | 3.937 | 11.529 -20.359 | 1.00 | 0.70 |
| MOTA | 433 | OE1 GLU | 32 | 4.969 | 12.161 -20.513 | | 1.37 |
| MOTA | 434 | OE2 GLU | 32 | 2.929 | 11.696 -21.026 | _ | 1.45 |
| ATOM | 435 | C GLU | 32 | | | | |
| | | | | 4.962 | 6.773 -19.235 | | 0.20 |
| MOTA | 436 | O GLU | 32 | 4.126 | 5.893 -19.280 | 1.00 | 0.20 |
| MOTA | 437 | n lys | 33 | 6.168 | 6.575 -19.689 | 1.00 | 0.20 |
| MOTA | 438 | hn lys | 33 | 6.835 | 7.293 -19.654 | | 0.21 |
| MOTA | 439 | CA LYS | 33 | 6.518 | 5.249 -20.269 | | 0.21 |
| ATOM | 440 | HA LYS | 33 | 5.825 | | | |
| MOTA | 441 | | | | 5.029 -21.068 | | 0.24 |
| | | CB LYS | 33 | 7.940 | 5.281 - 20.843 | 1.00 | 0.26 |
| ATOM | 442 | HB1 LYS | 33 | 7.987 | 6.024 -21.624 | 1.00 | 0.31 |
| MOTA | 443 | HB2 LYS | 33 | 8.179 | 4.312 -21.257 | | 0.31 |
| ATOM | 444 | CG LYS | 33 | 8.954 | 5.631 -19.748 | | |
| ATOM | 445 | HG1 LYS | 33 | | | | 0.26 |
| ATOM | 446 | | | 8.823 | 4.970 -18.906 | | 0.40 |
| | | HG2 LYS | 33 | 8.799 | 6.648 - 19.430 | 1.00 | 0.42 |
| MOTA | 447 | CD LYS | 33 | 10.380 | 5.469 -20.291 | 1.00 | 0.48 |
| ATOM | 448 | HD1 LYS | 33 | 10.466 | 4.517 -20.793 | | 0.74 |
| ATOM | 449 | HD2 LYS | 33 | 11.080 | 5.505 -19.469 | | 1.11 |
| ATOM | 450 | CE LYS | 33 | 10.705 | | | |
| MOTA | 451 | | | | 6.593 -21.282 | | 0.92 |
| | | HE1 LYS | 33 | 10.398 | 7.543 -20.868 | 1.00 | 1.52 |
| MOTA | 452 | HE2 LYS | 33 | 10.184 | 6.419 -22.211 | 1.00 | 1.19 |
| MOTA | 453 | NZ LYS | 33 | 12.172 | 6.614 -21.538 | | 1.60 |
| ATOM | 454 | HZ1 LYS | 33 | 12.668 | 6.957 -20.692 | | |
| MOTA | 455 | HZ2 LYS | 33 | | | | 1.99 |
| MOTA | | | | 12.374 | 7.247 - 22.340 | 1.00 | 2.14 |
| | 456 | HZ3 LYS | 33 | 12.498 | 5.653 -21.763 | 1.00 | 2.03 |
| MOTA | 457 | C LYS | 33 | 6.399 | 4.158 -19.202 | | 0.19 |
| MOTA | 458 | O LYS | 33 | 6.054 | 3.035 -19.495 | | 0.20 |
| MOTA | 459 | N ALA | 34 | 6.682 | | | |
| ATOM | 460 | | | | 4.471 -17.966 | | 0.17 |
| | | | 34 | 6.965 | 5.383 -17.740 | 1.00 | 0.18 |
| ATOM | 461 | CA ALA | 34 | 6.589 | 3.428 -16.904 | 1.00 | 0.16 |
| MOTA | 462 | HA ALA | 34 | 7.276 | 2.625 -17.128 | | 0.18 |
| ATOM | 463 | CB ALA | 34 | 6.952 | 4.043 -15.551 | | 0.16 |
| MOTA | 464 | HB1 ALA | 34 | 6.483 | 3.476 -14.761 | | |
| ATOM | 465 | HB2 ALA | 34 | | | | 1.02 |
| ATOM | 466 | | | 6.604 | 5.065 -15.516 | | 0.98 |
| | | | 34 | 8.024 | 4.022 - 15.423 | 1.00 | 1.02 |
| ATOM | 467 | C ALA | 34 | 5.164 | 2.875 -16.844 | 1.00 | 0.16 |
| MOTA | 468 | O ALA | 34 | 4.954 | 1.677 -16.847 | 1.00 | 0.17 |
| ATOM | . 469 | N PHE | 35 | 4.182 | 3.729 -16.792 | | 0.16 |
| ATOM | 470 | HN PHE | 35 | 4.364 | 4.694 -16.792 | | |
| MOTA | 471 | CA PHE | 35 | 2.781 | 3.230 -16.736 | | 0.16 |
| ATOM | 472 | HA PHE | | | | | 0.17 |
| ATOM | | | 35 | 2.690 | 2.525 -15.924 | 1.00 | 0.17 |
| | 473 | CB PHE | 35 | 1.815 | 4.396 -16.508 | 1.00 | 0.18 |
| ATOM | 474 | HB1 PHE | 35 | 0.802 | 4.060 -16.672 | | 0.19 |
| MOTA | 475 | HB2 PHE | 35 | 2.045 | 5.192 -17.200 | | 0.19 |
| ATOM | 476 | CG PHE | 35 | 1.953 | 4.902 -15.089 | | |
| MOTA | 477 | CD1 PHE | 35 | | | | 0.18 |
| MOTA | 478 | HD1 PHE | | 1.616 | 4.071 -14.011 | | 0.19 |
| | | | 35 | 1.258 | 3.069 -14.191 | 1.00 | 0.19 |
| MOTA | 479 | CD2 PHE | 35 | 2.415 | 6.203 -14.849 | 1.00 | 0.20 |
| MOTA | 480 | HD2 PHE | 35 | 2.674 | 6.847 -15.677 | | 0.21 |
| ATOM | 481 | CE1 PHE | 35 | 1.743 | 4.539 -12.699 | | 0.21 |
| ATOM | 482 | HE1 PHE | 35 | 1.484 | 3.897 -11.870 | | |
| MOTA | 483 | CE2 PHE | 35 | | | | 0.23 |
| ATOM | | | | 2.540 | 6.670 -13.535 | | 0.22 |
| | 484 | HE2 PHE | 35 | 2.893 | 7.672 -13.349 | 1.00 | 0.24 |
| MOTA | 485 | CZ PHE | 35 | 2.205 | 5.838 -12.460 | | 0.22 |
| MOTA | 486 | HZ PHE | 35 | 2.303 | 6.198 -11.447 | | 0.24 |
| MOTA | 487 | C PHE | 35 | 2.432 | | | |
| MOTA | 488 | O PHE | 35 | | 2.524 -18.048 | | 0.18 |
| ATOM | 489 | | | 1.770 | 1.507 -18.055 | | 0.19 |
| | | N LYS | 36 | 2.864 | 3.053 -19.162 | 1.00 | 0.19 |
| ATOM | 490 | HN LYS | 36 | 3.394 | 3.878 -19.144 | | 0.19 |
| MOTA | 491 | CA LYS | 36 | 2.535 | 2.399 -20.460 | | 0.22 |
| ATOM | 492 | HA LYS | 36 | 1.462 | 2.358 -20.574 | | |
| MOTA | 493 | CB LYS | 36 | | | | 0.23 |
| MOTA | 494 | | | 3.135 | 3.205 -21.614 | | 0.24 |
| | | HB1 LYS | 36 | 3.045 | 2.641 - 22.530 | | 0.27 |
| MOTA | 495 | HB2 LYS | 36 | 4.178 | 3.400 -21.412 | 1.00 | 0.24 |
| ATOM | 496 | CG LYS | 36 | 2.384 | 4.530 -21.758 | | 0.27 |
| MOTA | 497 | HG1 LYS | 36 | 2.471 | 5.097 -20.844 | | 0.69 |
| MOTA | 498 | HG2 LYS | 36 | 1.341 | 4.332 -21.963 | | |
| MOTA | 499 | CD LYS | 36 | 2.988 | 5.332 -22.913 | | 0.68 |
| ATOM | 500 | HD1 LYS | 36 | | A 766 00 000 | | 0.75 |
| MOTA | 501 | HD2 LYS | | 2.898 | 4.766 -23.828 | | 1.39 |
| | 201 | INC DID | 36 | 4.032 | 5.525 -22.710 | 1.00 | 1.34 |

| 10014 | E 0.0 | | | | | | | • |
|-------|------------|-------|-----|--------------|--------|---------------|----------|------|
| ATOM | 502 | CE | LYS | 36 | 2.243 | 6.659 -23. | 065 1.00 | 1.15 |
| MOTA | 503 | HE1 | LYS | 36 | 2.728 | 7.415 -22. | | |
| ATOM | 504 | HE2 | LYS | 36 | 1.221 | _ | | |
| MOTA | 505 | | LYS | 36 | | | | |
| ATOM | 506 | | • | | 2.260 | 7.076 -24. | | 1.99 |
| | | | LYS | 36 | 2.628 | 6.298 -25. | 079 1.00 | 2.51 |
| MOTA | 507 | HZ2 | LYS | 36 | 2.871 | 7.911 -24. | | 2.40 |
| ATOM | 508 | HZ3 | LYS | 36 | 1.295 | 7.309 -24. | | |
| ATOM | 509 | C | LYS | 36 | 3.098 | | | |
| ATOM | 510 | | LYS | | | 0.976 -20. | | 0.21 |
| | | | | 36 | 2.446 | 0.053 - 20. | | 0.23 |
| ATOM | 511 | | LYS | 37 | 4.295 | 0.778 - 19. | 995 1.00 | 0.21 |
| MOTA | 512 | HN | LYS | 37 | 4.810 | 1.527 - 19. | | 0.20 |
| MOTA | 513 | CA | LYS | 37 | 4.864 | -0.600 -19. | | |
| ATOM | 514 | | LYS | 37 | | | | 0.22 |
| MOTA | 515 | | | | 4.926 | -0.974 - 21. | | 0.24 |
| | | | LYS | 37 | 6.257 | -0.581 -19. | 358 1.00 | 0.22 |
| ATOM | 516 | | LYS | 37 | 6.589 | -1.596 -19. | | 0.24 |
| ATOM | 517 | HB2 | LYS | 37 | 6.216 | -0.061 -18. | | |
| ATOM | 518 | CG | LYS | 37 | 7.244 | 0.130 -20. | | 0.21 |
| MOTA | 519 | | LYS | 37 | | | | 0.26 |
| ATOM | 520 | | | | 6.921 | 1.140 -20. | | 0.25 |
| | | HG2 | | 37 | 7.296 | -0.398 -21.3 | 227 1.00 | 0.28 |
| MOTA | 521 | | LYS | 37 | 8.625 | 0.139 - 19.6 | | 0.30 |
| MOTA | 522 | HD1 | LYS | 37 | 8.994 | -0.873 - 19. | | |
| ATOM | 523 | HD2 | LYS | 37 | 8.549 | | | 0.77 |
| ATOM | 524 | | LYS | 37 | | | | 0.84 |
| ATOM | 525 | | LYS | | 9.594 | 0.968 -20. | | 0.90 |
| ATOM | | | | 37 | 10.530 | 1.076 -19. | | 1.47 |
| | 526 | | LYS | 37 | 9.169 | 1.945 -20. | 652 1.00 | 1.59 |
| MOTA | 527 | | LYS | 37 | 9.836 | 0.286 -21. | | 1.77 |
| MOTA | 528 | HZ1 | LYS | 37 | 9.798 | 0.984 -22. | | |
| MOTA | 529 | HZ2 | | 37 | 9.106 | _ | | 2.22 |
| ATOM | 530 | | LYS | 37 | | -0.439 -21. | | 2.28 |
| MOTA | 531 | | | | 10.774 | -0.161 -21. | | 2.33 |
| | | | LYS | 37 | 3.955 | -1.506 -19.3 | 158 1.00 | 0.20 |
| ATOM | 532 | | LYS | 37 | 3.689 | -2.636 -19. | 516 1.00 | 0.21 |
| MOTA | 533 | N | ALA | 38 | 3.479 | -1.013 -18.0 | | 0.19 |
| ATOM | 534 | HN | ALA | 38 | 3.711 | -0.098 -17. | | |
| ATOM | 53\$ | CA · | ALA | 38 | 2.589 | -1.838 -17. | | 0.19 |
| MOTA | 536 | HA . | ALA | 38 | 3.116 | | | 0.18 |
| ATOM | 537 | | ALA | 38 | | -2.727 -16. | | 0.19 |
| ATOM | 538 | | ALA | | 2.183 | -1.030 -15.9 | | 0.19 |
| MOTA | 539 | | | 38 | 2.831 | -0.172 -15.8 | | 1.05 |
| | _ | HB2 | | 38 | 2.270 | -1.649 - 15.0 | 068 1.00 | 1.00 |
| ATOM | 540 | | ALA | 38 | 1.161 | -0.698 -16.0 | | 1.06 |
| MOTA | 541 | C. | ALA | 38 | 1.338 | -2.238 -17.9 | | |
| MOTA | 542 | 0 | ALA | 38 | 0.967 | -3.392 -18.0 | • • • | 0.18 |
| MOTA | 543 | N | PHE | 39 | 0:688 | | | 0.19 |
| MOTA | 544 | | PHE | 39 · | | -1.295 -18.9 | | 0.18 |
| MOTA | 545 | _ | PHE | | 1.005 | -0.368 -18. | | 0.18 |
| ATOM | 546 | | | 39 | -0.535 | -1.632 - 19.3 | | 0.19 |
| ATOM | | | PHE | 39 | -1.248 | -2.122 -18.7 | 720 1.00 | 0.19 |
| | 547 | | PHE | 3 9 · | -1.156 | -0.354 - 19.9 | | 0.21 |
| MOTA | 548 | HB1 | PHE | 39 | -1.883 | -0.614 - 20.6 | | 0.24 |
| MOTA | 549 | HB2 | PHE | 39 | -0.381 | 0.256 -20.3 | | |
| ATOM | 550 | CG | PHE | 39 | -1.836 | | | 0.21 |
| MOTA | 551 | | PHE | 39 | | 0.416 -18.8 | | 0.20 |
| ATOM | 552 | | PHE | | -3.010 | -0.080 -18.2 | | 0.25 |
| ATOM | 553 | | | 39 | -3.429 | -1.014 - 18.5 | | 0.30 |
| | | | PHE | 39 | -1.294 | 1.627 -18.3 | 380 1.00 | 0.17 |
| ATOM | 554 | | PHE | 39 | -0.389 | 2.012 -18.8 | | 0.18 |
| ATOM | 555 | CE1 | PHE | 39 | -3.642 | 0.633 -17.2 | | 0.28 |
| ATOM | 556 | HE1 | PHE | 39 | -4.548 | 0.250 -16.7 | | |
| MOTA | 557 | CE2 | PHE | 39 | -1.926 | | | 0.34 |
| MOTA | 558 | | PHE | 39 | | 2.341 -17.3 | | 0.18 |
| ATOM | 559 | | PHE | | -1.507 | 3.275 -17.0 | | 0.17 |
| ATOM | 560 | | | 39 | -3.099 | 1.843 -16.7 | | 0.23 |
| ATOM | | | PHE | 39 | -3.587 | 2.394 -15.5 | 85 1.00 | 0.26 |
| | 561 | | PHE | 39 | -0.154 | -2.571 - 20.5 | 08 1.00 | 0.18 |
| MOTA | 562 | 0 | PHE | 39 | -0.862 | -3.509 -20.8 | | |
| MOTA | 563 | N I | LYS | 40 | 0.963 | -2.330 -21.1 | | 0.18 |
| ATOM | 564 | HN 1 | LYS | 40 | 1.522 | | | 0.19 |
| MOTA | 565 | | LYS | 40 | | -1.570 -20.8 | | 0.19 |
| MOTA | 566 | | LYS | | 1.388 | -3.214 -22.2 | | 0.19 |
| ATOM | 567 | | | 40 | 0.642 | -3.186 -23.0 | 31 1.00 | 0.20 |
| | | | LYS | 40 | 2.730 | -2.707 - 22.8 | 04 1.00 | 0.21 |
| MOTA | 568 | HB1 I | | 40 | 3.466 | -2.723 -22.0 | | 0.21 |
| ATOM | 569 | HB2 I | | 40 | 2.610 | -1.692 -23.1 | | |
| ATOM | 570 | CG I | LYS | 40 | 3.218 | -3.588 -23.9 | | 0.25 |
| ATOM | 571 | HG1 I | | 40 | 3.337 | | | 0.25 |
| ATOM | 572 | HG2 I | | 40 | 4.171 | -4.604 -23.6 | | 0.46 |
| MOTA | 573 | | LYS | 40 | | -3.218 -24.3 | | 0.46 |
| ATOM | 574 | HD1 I | | | 2.213 | -3.560 -25.1 | 21 1.00 | 0.38 |
| MOTA | 575 | | | 40 | 1.840 | -2.555 -25.2 | 53 1.00 | 0.54 |
| ATOM | 576 | HD2 I | | 40 | 1.392 | -4.227 -24.9 | 05 1.00 | 0.56 |
| ATOM | | | LYS | 40 | 2.903 | -4.019 - 26.4 | 07 1.00 | 0.40 |
| | 577 570 | HE1 I | | 40 | 3.776 | -4.604 -26.1 | | 1.07 |
| MOTA | 578 | HE2 I | LYS | 40 | 3 100 | -3 157 -26 0 | | 1.07 |
| | | | | | | | | |

| ATOM | 579 NZ LYS | 40 | 1.958 | -4.852 -27.203 | 1 00 1 10 |
|--------------|------------------------------|----------|------------------|-------------------------------------|------------------------|
| ATOM | 580 HZ1 LYS | 40 | 1.571 | -5.607 -26.602 | 1.00 1.40 1.00 1.95 |
| MOTA | 581 HZ2 LYS | | 2.464 | -5.274 -28.009 | 1.00 1.92 |
| ATOM | 582 HZ3 LYS | 40 | 1.181 | -4.258 -27.552 | 1.00 2.02 |
| MOTA MOTA | 583 C LYS 584 O LYS | 40 | 1.553 | -4.648 - 21.740 | 1.00 0.17 |
| MOTA | 584 O LYS 585 N VAL | 40 | 1.034 | -5.583 -22.314 | 1.00 0.17 |
| ATOM | 586 HN VAL | 41 | 2.271 2.681 | -4.828 - 20.663 | 1.00 0.17 |
| MOTA | 587 CA VAL | 41 | 2.468 | -4.060 -20.214 -6.204 -20.116 | 1.00 0.18 |
| MOTA | 588 HA VAL | | 2.953 | -6.816 -20.862 | 1.00 0.16 1.00 0.17 |
| ATOM | 589 CB VAL | 41 | 3.350 | -6.143 -18.868 | 1.00 0.18 |
| ATOM | 590 HB VAL | 41 | 2.966 | -5.393 -18.192 | 1.00 0.41 |
| MOTA MOTA | 591 CG1 VAL 592 HG11 VAL | 41 | 3.343 | -7.508 -18.175 | 1.00 0.44 |
| MOTA | 592 HG11 VAL 593 HG12 VAL | 41 | 2.420 | -7.631 -17.629 | 1.00 1.16 |
| ATOM | 594 HG13 VAL | 41 41 | 4.176 3.429 | -7.571 -17.490 -8.289 -18.916 | 1.00 1.18 |
| ATOM | 595 CG2 VAL | | 4.781 | -8.289 -18.916 -5.785 -19.277 | 1.00 1.11 1.00 0.43 |
| MOTA | 596 HG21 VAL | 41 | 5.132 | -6.492 -20.013 | 1.00 0.43 1.00 1.12 |
| MOTA | 597 HG22 VAL | | 5.423 | -5.820 -18.411 | 1.00 1.11 |
| MOTA | 598 HG23 VAL | 41 | 4.797 | -4.790 -19.697 | 1.00 1.19 |
| MOTA MOTA | 599 C VAL 600 O VAL | 41 | 1.122 | -6.833 - 19.751 | 1.00 0.16 |
| MOTA | 600 O VAL 601 N TRP | 41 | 0.887 | -7.999 -19.996 | 1.00 0.17 |
| ATOM | 602 HN TRP | | 0.240 0.448 | -6.080 -19.152 -5.143 -18.950 | 1.00 0.16 |
| MOTA | 603 CA TRP | 42 | -1.079 | -6.655 -18.761 | 1.00 0.17 1.00 0.17 |
| ATOM | 604 HA TRP | 42 | -0.927 | -7.642 -18.352 | 1.00 0.17 |
| MOTA | 605 CB TRP | | -1.739 | -5.767 -17.699 | 1.00 0.18 |
| MOTA | 606 HB1 TRP | 42 | -2.787 | -6.018 -17.621 | 1.00 0.19 |
| MOTA MOTA | 607 HB2 TRP 608 CG TRP | 42 | -1.638 | -4.730 - 17.983 | 1.00 0.20 |
| MOTA | 608 CG TRP 609 CD1 TRP | 42 42 | -1.073 | -5.990 -16.377 | 1.00 0.18 |
| ATOM | 610 HD1 TRP | 42 | -0.311 -0.092 | -5.082 -15.724 | 1.00 0.22 |
| MOTA | 611 CD2 TRP | 42 | -1.095 | -4.084 -16.066 -7.182 -15.539 | 1.00 0.28 1.00 0.19 |
| MOTA | 612 NE1 TRP | 42 | 0.140 | -5.643 -14.543 | 1.00 0.19 1.00 0.22 |
| ATOM | 613 HE1 TRP | 42 | 0.714 | -5.194 -13.887 | 1.00 0.25 |
| MOTA MOTA | 614 CE2 TRP | 42 | -0.315 | -6.935 - 14.384 | 1.00 0.20 |
| MOTA | 615 CE3 TRP 616 HE3 TRP | 42 | -1.707 | -8.441 -15.669 | 1.00 0.25 |
| ATOM | 617 CZ2 TRP | 42 42 | -2:309 -0:149 | -8.658 -16.539 | 1.00 0.27 |
| ATOM | 618 HZ2 TRP | 42 | 0.454 | -7.903 -13.393 -7.691 -12.521 | 1.00 0.24 |
| MOTA | 619 CZ3 TRP | 42 | -1.543 | -9.418 -14.673 | 1.00 0.25 1.00 0.31 |
| MOTA | 620 HZ3 TRP | 42 | | -10.381 -14.782 | 1.00 0.31 |
| MOTA | 621 CH2 TRP | 42 | -0.764 | -9.149 -13.538 | 1.00 0.30 |
| ATOM ATOM | 622 HH2 TRP 623 C TRP | 42 | -0.642 | -9.904 - 12.775 | 1.00 0.35 |
| MOTA | 623 C TRP 624 O TRP | 42 | -1.991 | -6.754 -19.985 | 1.00 0.17 |
| MOTA | 625 N SER | 42 43 | -2.726 -1.952 | -7.706 -20.138 -5.702 -20.055 | 1.00 0.18 |
| MOTA | 626 HN SER | 43 | ~1.352 | -5.782 -20.855 -5.021 -20.713 | 1.00 0.17 |
| MOTA | 627 CA SER | 43 | -2.831 | -5.825 -22.062 | 1.00 0.17 1.00 0.18 |
| ATOM | 628 HA SER | 43 | -3.846 | -6.028 -21.759 | 1.00 0.19 |
| MOTA | 629 CB SER | 43 | -2.779 | -4.474 -22.775 | 1.00 0.20 |
| MOTA MOTA | 630 HB1 SER 631 HB2 SER | 43 | -2.965 | -3.683 - 22.059 | 1.00 0.21 |
| MOTA | 631 HB2 SER 632 OG SER | 43 43 | -3.533 -1.400 | -4.442 -23.543 | 1.00 0.23 |
| ATOM | 633 HG SER | 43 | -1.499 -1.031 | -4.304 -23.368 -5.140 -23.309 | 1.00 0.21 |
| MOTA | 634 C SER | 43 | -2.358 | -6.922 -23.019 | 1.00 0.97 1.00 0.18 |
| MOTA | 635 O SER | 43 | -3.085 | -7.350 -23.893 | 1.00 0.21 |
| ATOM ATOM | 636 N ASP | 44 | -1.148 | -7.379 -22.866 | 1.00 0.17 |
| ATOM | 637 HN ASP 638 CA ASP | 44 | -0.575 | -7.019 -22.156 | 1.00 0.18 |
| ATOM | 639 HA ASP | 44 44 | -0.632 -0.650 | -8.445 -23.770 | 1.00 0.18 |
| ATOM | 640 CB ASP | 44 | -0.650 0.809 | -8.086 -24.788 -8.793 -23.386 | 1.00 0.19 |
| MOTA | 641 HB1 ASP | .44 | 1.117 | -9.683 -23.915 | 1.00 0.20 1.00 0.21 |
| ATOM | 642 HB2 ASP | 44 | 0.864 | -8.969 -22.322 | 1.00 0.21 1.00 0.22 |
| ATOM | 643 CG ASP | 44 | 1.734 | -7.635 -23.760 | 1.00 0.24 |
| MOTA MOTA | 644 OD1 ASP 645 OD2 ASP | 44 | 1.340 | -6.833 - 24.591 | 1.00 0.85 |
| ATOM | 645 OD2 ASP 646 C ASP | 44 44 | | -7.568 -23.209 | 1.00 0.84 |
| ATOM | 647 O ASP | 44 | -1.499 -1.753 | -9.705 -23.665 -10.366 -24.653 | 1.00 0.19 |
| MOTA | 648 N VAL | 45 | -1.927 | -10.366 - 24.653 $-10.058 - 22.475$ | 1.00 0.21 |
| MOTA | 649 HN VAL | 45 | -1.689 | -9.519 -21.693 | 1.00 0.21 1.00 0.21 |
| MOTA | 650 CA VAL | 45 | -2.749 | -11.299 -22.302 | 1.00 0.21 |
| MOTA MOTA | 651 HA VAL | 45 | -2.833 · | -11.811 -23.247 | 1.00 0.28 |
| ATOM | 652 CB VAL 653 HB VAL | 45 45 | -2.045 | -12.222 -21.303 | 1.00 0.30 |
| ATOM | 653 HB VAL 654 CG1 VAL | 45 45 | -2.645 · | -13.107 -21.146 | 1.00 0.37 |
| MOTA | 655 HG11 VAL | 45 | -0.076 | -12.626 -21.866 -11 766 -22 323 | 1.00 0.36 |
| | | | | | |

| MOTA | 656 HG12 VAI | 45 | -0.810 | -13.400 | -22 607 | 1.00 | 1.02 |
|-----------------|------------------------------|------|--------------------|---------|--------------------|--------------|--------------|
| ATOM | 657 HG13 VAI | | | -12.995 | | 1.00 | 1.13 |
| MOTA | 658 CG2 VAI | | | -11.486 | | 1.00 | 0.32 |
| MOTA | 659 HG21 VAI | | | -11.303 | -19.524 | 1.00 | 0.96 |
| MOTA | 660 HG22 VAI | 45 | | -10.545 | | 1.00 | 1.09 |
| MOTA | 661 HG23 VAI | 45 | -1.258 | -12.091 | | 1.00 | 1.11 |
| ATOM | 662 C VAI | 45 | -4.160 | -10.966 | -21.790 | 1.00 | 0.29 |
| MOTA | 663 O VAI | 45 | -4.837 | -11.819 | -21.249 | 1.00 | 0.64 |
| MOTA | 664 N THE | 46 | -4.619 | -9.748 | -21.963 | 1.00 | 0.36 |
| MOTA | 665 HN THE | | -4.062 | -9.076 | -22.409 | 1.00 | 0.65 |
| MOTA | 666 CA THE | | -5.998 | -9.382 | -21.491 | 1.00 | 0.38 |
| MOTA | 667 HA THE | | | | -21.320 | 1.00 | 0.44 |
| ATOM | 668 CB THE | | -5.912 | | -20.186 | 1.00 | 0.39 |
| MOTA | 669 HB TH | | -6.889 | -8.193 | -19.943 | 1.00 | 0.46 |
| ATOM | 670 OG1 THE | | -5.018 | | -20.358 | 1.00 | 0.36 |
| ATOM | 671 HG1 TH | | -5.532 | | -20.608 | 1.00 | 0.94 |
| MOTA MOTA | 672 CG2 THI | | -5.430 | | -19.036 | 1.00 | 0.43 |
| MOTA | 673 HG21 THE 674 HG22 THE | | -4.929 | -10.327 | | 1.00 | 1.08 |
| MOTA | | | -6.277 | | -18.445 | 1.00 | 1.15 |
| MOTA | 675 HG23 THI 676 C THI | | -4.746 | | -18.415 | 1.00 | 1.05 |
| ATOM | 677 O TH | | -6.668 -6.124 | | -22.553 | 1.00 | 0.32 |
| ATOM | 678 N PRO | | -7.833 | | -22.892 | 1.00 | 0.32 |
| ATOM | 679 CA PRO | | -8.479 | | -23.084 -24.100 | 1.00 | 0.30 |
| MOTA | 680 HA PRO | | -7.820 | | -24.100 | 1.00 | 0.30 |
| MOTA | 681 CB PRO | | -9.687 | | -24.546 | 1.00 | 0.33 |
| MOTA | 682 HB1 PR | | -9.541 | | -25.561 | | 0.35 |
| MOTA | 683 HB2 PRO | | -10.579 | | -24.489 | 1.00 | 0.40 |
| MOTA | 684 CG PR | | -9.825 | | -23.621 | 1.00 | 0.37 |
| MOTA | 685 HG1 PRO | | | -10.885 | | 1.00 | 0.35 |
| MOTA | 686 HG2 PR | | -10.703 | | -23.001 | 1.00 | 0.34 |
| MOTA | 687 CD PRO | | | -10.077 | | 1.00 | 0.33 |
| MOTA | 688 HD2 PR | | -8.853 | | -21.692 | 1.00 | 0.31 |
| MOTA | 689 HD1 PR | | | -10.946 | | 1.00 | 0.39 |
| MOTA | 690 C PR | | -8.933 | | -23.506 | 1.00 | 0.25 |
| MOTA | 691 O PR | 47 | -9.744 | | -24.080 | 1.00 | |
| MOTA | 692 N LE | | -8.418 | | -22.362 | 1.00 | 0.26 |
| MOTA | 693 HN LE | J 48 | -7.766 | | -21.912 | 1.00 | 0.29 |
| MOTA | 694 CA LE | J 48 | -8.827 | | -21.742 | 1.00 | 0.26 |
| MOTA | 695 HA LE | J 48 | -9.904 | | -21.696 | 1.00 | 0.27 |
| MOTA | 696 CB LE | J 48 | -8.241 | | -20.329 | 1.00 | 0.31 |
| ATOM | 697 HB1 LE | | -8.476 | -3.892 | -19.909 | 1.00 | |
| MOTA | 698 HB2 LE | | -7.167 | | -20.385 | 1.00 | 0.33 |
| ATOM | 699 CG LE | | -8.816 | -5.964 | -19.434 | 1.00 | 0.34 |
| MOTA | 700 HG LE | - | -8.808 | | -19.972 | 1.00 | 0.32 |
| ATOM | 701 CD1 LE | | -7.952 | | -18.177 | 1.00 | 0.41 |
| MOTA | 702 HD11 LE | | -8.002 | | -17.613 | 1.00 | 1.11 |
| MOTA MOTA | 703 HD12 LE | | -6.928 | | -18.462 | 1.00 | 1.05 |
| ATOM | 704 HD13 LE 705 CD2 LE | _ | -8.315 | | -17.570 | 1.00 | 1.15 |
| ATOM | 705 CD2 LE 706 HD21 LE | | -10.255 | | -19.016 | 1.00 | 0.36 |
| ATOM | 707 HD22 LE | | -10.569 | | -19.478 | 1.00 | 1.10 |
| MOTA | 708 HD23 LE | _ | -10.299 | _ | -17.942 | 1.00 | 1.09 |
| ATOM | 709 C LE | | -10.912 -8.289 | | -19.325 | 1.00 | 1.04 |
| MOTA | 710 O LE | | -7.174 | | -22.589 -23.071 | 1.00 | 0.25 |
| MOTA | 711 N AS | | -9.073 | | -22:762 | 1.00 | 0.26 |
| ATOM | 712 HN AS | | -9.964 | | -22.762 | 1.00 1.00 | 0.25 0.26 |
| MOTA | 713 CA AS | | -8.622 | | -23.568 | 1.00 | 0.25 |
| MOTA | 714 HA AS | | -7.703 | | -24.082 | 1.00 | 0.27 |
| ATOM | 715 CB AS | | -9.700 | | -24.593 | 1.00 | 0.28 |
| ATOM | 716 HB1 AS | | -9.390 | | -25.153 | 1.00 | 0.30 |
| MOTA | 717 HB2 AS | | -10.628 | | -24.081 | 1.00 | 0.28 |
| MOTA | 718 CG AS | N 49 | -9.902 | | -25.553 | 1.00 | 0.32 |
| MOTA | 719 OD1 AS | N 49 | -9.798 | | -25.161 | 1.00 | 1.10 |
| MOTA | 720 ND2 AS | | -10.186 | | -26.804 | 1.00 | 1.14 |
| MOTA | 721 HD21 AS | | -10.268 | | -27.121 | 1.00 | 1.94 |
| MOTA | 722 HD22 AS | | -10.317 | -2.927 | -27.427 | 1.00 | 1.14 |
| MOTA | 723 C AS | | -8.391 | -0.417 | -22.633 | 1.00 | 0.24 |
| MOTA | 724 O AS | | -9.290 | | -21.939 | 1.00 | 0.23 |
| MOTA | 725 N PH | | -7.192 | | -22.606 | 1.00 | 0.24 |
| ATOM | 726 HN PH | | -6.485 | | -23.173 | 1.00 | 0.26 |
| MOTA | 727 CA PH | | -6.896 | | -21.710 | 1.00 | 0.23 |
| ATOM | 728 HA PH | | -7.688 | | -20.985 | 1.00 | 0.21 |
| MOTA MOTA | 729 CB PH | | -5.574 | | -20.981 | 1.00 | 0.24 |
| ATOM | 730 HB1 PH 731 HB2 PH | | - 5.357 | | -20.334 | 1.00 | 0.25 |
| ATOM | 731 NB2 PH | | -4.780 -5.676 | | -21.705 -20 154 | 1.00 | 0.27 |
| V 44 | , J. L.G FA | | -7 R/R | /4 (| -/// 15/ | חת ד | U JĄ |

| ATOM | 733 | CD1 | PHE | 50 | -6.266 | -0.201 - | 10 006 | 1.00 | 0.25 |
|------|-----|----------------|------|-----------|-----------------|----------|---------|------|------|
| ATOM | 734 | HD1 | PHE | 50 | -6.652 | | | | |
| ATOM | 735 | CD2 | | | | 0.731 - | | 1.00 | 0.28 |
| | | | PHE | 50 | -5.176 | -1.451 - | 20.654 | 1.00 | 0.22 |
| MOTA | 736 | HD2 | PHE | 50 | -4.720 | -1.483 - | -21.633 | 1.00 | 0.23 |
| ATOM | 737 | ÇE1 | PHE | 50 | -6.358 | -1.368 - | -18.117 | 1.00 | 0.25 |
| MOTA | 738 | HE1 | PHE | 50 | -6.813 | -1.336 - | | 1.00 | |
| MOTA | 739 | CE2 | PHE | 50 | | | • | | 0.28 |
| • | | _ | | | -5.267 | -2.618 - | | 1.00 | 0.23 |
| ATOM | 740 | HE2 | PHE | 50 | -4.881 | -3.550 - | -20.272 | 1.00 | 0.25 |
| ATOM | 741 | CZ | PHE | 50 | -5.858 | -2.576 - | 18.618 | 1.00 | 0.24 |
| ATOM | 742 | HZ | PHE | 50 | -5.928 | -3.476 - | | 1.00 | 0.25 |
| ATOM | 743 | C | PHE | 50 | -6.777 | | _ | | |
| ATOM | 744 | ŏ | | | | 2.538 - | | 1.00 | 0.26 |
| | | - | PHE | 50 | -6.028 | 2.596 - | | 1.00 | 0.31 |
| MOTA | 745 | N | THR | 51 | -7.517 | 3.555 - | -22.184 | 1.00 | 0.24 |
| ATOM | 746 | HN | THR | 51 | -8.109 | 3.468 - | -21.413 | 1.00 | 0.22 |
| MOTA | 747 | CA | THR | 51 | -7.470 | 4.842 - | | 1.00 | 0.27 |
| ATOM | 748 | HA | THR | 51 | -6.775 | | | | |
| ATOM | 749 | CB | THR | | | 4.762 - | | 1.00 | 0.31 |
| | | | | 51 | -8.868 | 5.153 - | | 1.00 | 0.30 |
| ATOM | 750 | | THR | 51 | -9.562 | 5.248 - | -22.663 | 1.00 | 0.29 |
| MOTA | 751 | OG1 | THR | 51 | -9.283 | 4.100 - | -24.341 | 1.00 | 0.35 |
| MOTA | 752 | HG1 | THR | 51 | -9.638 | 4.491 - | | 1.00 | 0.84 |
| ATOM | 753 | CG2 | THR | 51 | -8.835 | | | | |
| ATOM | 754 | | | | | 6.464 - | _ | 1.00 | 0.34 |
| | | | THR | 51 | -9.805 | | -24.716 | 1.00 | 1.02 |
| ATOM | 755 | | THR | 51 | -8.092 | 6.394 - | -25.053 | 1.00 | 1.07 |
| ATOM | 756 | HG23 | THR | 51 | -8.588 | 7.280 - | -23.611 | 1.00 | 1.13 |
| MOTA | 757 | C | THR | 51 | -7.024 | 5.969 - | | 1.00 | 0.25 |
| ATOM | 758 | 0 | THR | 51 | -7.553 | | | - | |
| ATOM | 759 | N | ARG | | | | -20.920 | 1.00 | 0.22 |
| | | - | - | 52 | -6.054 | 6.740 - | | 1.00 | 0.29 |
| ATOM | 760 | HN | ARG | 52 | -5.645 | 6.583 - | -23.287 | 1.00 | 0.32 |
| ATOM | 761 | CA | ARG | 52 | -5.566 | 7.861 - | -21.556 | 1.00 | 0.29 |
| ATOM | 762 | HA | ARG | 52 | -5.591 | | -20.518 | 1.00 | 0.27 |
| MOTA | 763 | CB | ARG | 52 | | | | | |
| | | | | | -4.128 | | -21.955 | 1.00 | 0.35 |
| ATOM | 764 | | ARG | 52 | -4.125 | 8.654 - | -22.935 | 1.00 | 0.39 |
| ATOM | 765 | HB2 | ARG | 52 | -3.539 | 7.295 - | -21.977 | 1.00 | 0.38 |
| ATOM | 766 | CG | ARG | 52 | · -3.521 | | -20.945 | 1.00 | 0.39 |
| ATOM | 767 | HG1 | ARG | 52 | -3.645 | | 19.946 | | |
| ATOM | 768 | | ARG | 52 | | | | 1.00 | 0.71 |
| ATOM | | | | | -4.017 | 10.134 - | | 1.00 | 0.57 |
| | 769 | | ARG | 52 | -2.030 | 9.345 - | -21.244 | 1.00 | 0.79 |
| MOTA | 770 | | ARG | 52 | -1.825 | 9.001 - | -22.248 | 1.00 | 1.45 |
| ATOM | 771 | HD2 | ARG | 52 | -1.453 | | -20.543 | 1.00 | 1.39 |
| ATOM | 772 | NE | ARG | 52 | -1.656 | 10.782 - | _ | _ | |
| ATOM | 773 | HE | ARG | 52 | -2.354 | | | 1.00 | 1.47 |
| ATOM | 774 | CZ | ARG | | | 11.468 - | | 1.00 | 2.06 |
| | _ | | | 52 | -0.398 | 11.127 - | | 1.00 | 2.09 |
| ATOM | 775 | | | 52 | -0.070 | 12.385 - | -20.960 | 1.00 | 3.05 |
| ATOM | 776 | _ | _ | 52 | -0.782 | 13.084 - | -20.911 | 1.00 | 3.45 |
| ATOM | 777 | HH12 | ARG | 52 | 0.894 | 12.649 - | -20.923 | 1.00 | 3.60 |
| ATOM | 778 | NH2 | ARG | 52 | 0.532 | 10.213 - | | 1.00 | |
| ATOM | 779 | | | 52 | 0.281 | | _ | | 2.31 |
| ATOM | 780 | | | | | | -21.226 | 1.00 | 2.16 |
| | | | | 52 | 1.496 | 10.477 - | | 1.00 | 3.05 |
| ATOM | 781 | C | ARG | 52 | -6.460 | 9.090 - | -21.758 | 1.00 | 0.29 |
| MOTA | 782 | 0 | ARG | 52 | -6.719 | 9.495 - | -22.875 | 1.00 | 0.33 |
| MOTA | 783 | N | LEU | 53 | -6.928 | | -20.689 | 1.00 | 0.26 |
| MOTA | 784 | HN | LEU | 53 | -6.702 | | _ | | |
| ATOM | 785 | CA | LEU | | | | -19.798 | 1.00 | 0.25 |
| ATOM | | | | 53 | -7.803 | 10.896 - | | 1.00 | 0.29 |
| | 786 | HA | LEU | 53 | -8.167 | 10.972 - | -21.835 | 1.00 | 0.32 |
| MOTA | 787 | | LEU | 53 | -8.992 | 10.784 - | -19.862 | 1.00 | 0.28 |
| MOTA | 788 | HB1 | LEU | 53 | -9.579 | 11.688 - | | 1.00 | 0.31 |
| MOTA | 789 | HB2 | LEU | 53 | -8.624 | 10.648 - | | 1.00 | 0.28 |
| MOTA | 790 | CG | LEU | 53 | -9.866 | | | | - |
| ATOM | 791 | HG | LEU | | | | -20.249 | 1.00 | 0.28 |
| | | | _ | 53 | -9.264 | | -20.246 | 1.00 | 0.29 |
| MOTA | 792 | | LEU | 53 | -10.999 | 9.440 - | -19.232 | 1.00 | 0.29 |
| ATOM | | HD11 | | 53 | -11.606 | 8.585 - | -19.487 | 1.00 | 0.95 |
| MOTA | 794 | HD12 | LEU | 53 | -11.610 | 10.331 - | • | 1.00 | |
| ATOM | | HD13 | | 53 | -10.581 | | | | 1.05 |
| ATOM | 796 | | | | | | -18.247 | 1.00 | 1.07 |
| ATOM | | | | 53 | -10.463 | | -21.646 | 1.00 | 0.36 |
| | 797 | | | 53 | -10.523 | 10.856 - | | 1.00 | 1.01 |
| ATOM | 798 | | _ | 53 | -11.453 | 9.370 - | -21.685 | 1.00 | 1.09 |
| ATOM | 799 | HD23 | LEU | 53 | -9.835 | 9.319 - | | 1.00 | 1.14 |
| MOTA | 800 | | LEU | 53 | -7.000 | 12.154 | | | |
| ATOM | 801 | ŏ | LEU | 53 | | | | 1.00 | 0.33 |
| ATOM | 802 | _ | | | -6.315 | 12.218 - | | 1.00 | 0.34 |
| | | | HIS | 54 | -7.080 | 13.154 - | | 1.00 | 0.41 |
| MOTA | 803 | HN | HIS | 54 | -7.637 | 13.075 - | | 1.00 | 0.45 |
| ATOM | 804 | CA | HIS | 54 | -6.324 | 14.413 - | | 1.00 | 0.47 |
| MOTA | 805 | HA | HIS | 54 | -5.292 | 14.183 - | | 1.00 | 0.54 |
| MOTA | 806 | | HIS | 54 | -6.407 | 15.314 - | | | |
| ATOM | 807 | HB1 | | 54 | -6.018 | 16.291 | | 1.00 | 0.60 |
| MOTA | 808 | | HIS | 54 | -7.438 | | | 1.00 | 0.64 |
| ATOM | 809 | CG | HIS | | | 15.407 - | | 1.00 | 0.61 |
| | 003 | - G | 1172 | 54 | -5.602 | 14.726 - | -23.426 | 1.00 | 0.74 |

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| ATOM | 810 | ND1 | UTC | . 54 | -5.645 | 15 054 04 | 707 4 | | |
|-------|-----|------|-----|-----------------|---------|--------------|--------------|------|-------|
| | | | | | | 15.254 -24. | _ | 1.00 | 1.35 |
| MOTA | 811 | HD1 | HIS | 54 | -6.172 | 16.028 -24. | 996 1 | 1.00 | 1.86 |
| MOTA | 812 | CD2 | HIS | 54 | -4.740 | 13.656 -23. | | 1.00 | 0.86 |
| MOTA | 813 | HD2 | | 54 | | | | | |
| | | | | | -4.480 | 13.010 -22. | 008] | L.00 | 1.34 |
| ATOM | 814 | CE1 | HIS | 54 | -4.834 | 14.512 -25. | 481 1 | 1.00 | 1.33 |
| MOTA | 815 | HE1 | HIS | 54 | -4.670 | 14.692 -26. | | | |
| | | | | | | | | L.00 | 1.83 |
| MOTA | 816 | NE2 | HIS | 54 | -4.257 | 13.525 -24. | 792 1 | 1.00 | 0.92 |
| MOTA | 817 | C | HIS | 54 | -6.933 | 15.154 -19. | _ | | |
| | | | | | | | | L.00 | 0.43 |
| MOTA | 818 | 0 | HIS | 54 | -6.230 | 15.714 -19. | 051 1 | L.00 | 0.49 |
| MOTA | 819 | N | ASP | 55 | -8.236 | 15.172 -19. | | | _ |
| MOTA | | | | | | | | L.00 | 0.42 |
| | 820 | HN | ASP | 55 | -8.784 | 14.719 -20. | 442] | L.00 | 0.45 |
| ATOM. | 821 | CA | ASP | 55 | -8.892 | 15.892 -18. | 635 1 | 1.00 | 0.49 |
| MOTA | 822 | HA | ASP | 55 | _ | | | | _ |
| | | | | | -8.217 | 15.938 -17. | | 1.00 | 0.54 |
| MOTA | 823 | CB | ASP | 55 | -9.251 | 17.314 -19. | 073 1 | L.00 | 0.65 |
| MOTA | 824 | HB1 | ASP | [•] 55 | -9.876 | 17.774 -18. | | | |
| | | | | | | | | L.00 | 0.75 |
| ATOM | 825 | | asp | 55 | -9.783 | 17.277 -20. | 013 1 | 1.00 | 0.68 |
| ATOM | 826 | CG | ASP | 55 | -7.974 | 18.140 -19. | | 1.00 | |
| MOTA | 827 | OD1 | | | | | | | 0.71 |
| | | | | 55 ' | -7.978 | 19.037 -20. | 071] | 1.00 | 1.19 |
| ATOM | 828 | OD2 | ASP | 55 | -7.018 | 17.870 -18. | | 1.00 | 1.28 |
| MOTA | 829 | С | ASP | 55 | -10.167 | | | | |
| | | | | | | 15.156 -18. | | L.00 | 0.45 |
| MOTA | 830 | 0 | ASP | 55 | -10.638 | 14.273 -18. | 912 1 | 1.00 | 0.44 |
| MOTA | 831 | N | GLY | 56 | -10.728 | 15.518 -17. | | | |
| MOTA | 832 | | | | | | | L.00 | 0.46 |
| | | HN | GLY | 56 | -10.328 | 16.233 -16. | 563 | 1.00 | 0.50 |
| MOTA | 833 | CA | GLY | 56 | -11.975 | 14.848 -16. | | L.00 | 0.44 |
| MOTA | 834 | HA1 | GLY | 56 | -12.482 | | | | |
| | | | | | | 14.399 -17. | | 1.00 | 0.44 |
| MOTA | 835 | HA2 | GLY | 56 | -12.622 | 15.579 -16. | 169 1 | 1.00 | 0.48 |
| MOTA | 836 | C | GLY | 56 | -11.624 | | | | |
| | | | | | | 13.760 -15. | | 1.00 | 0.40 |
| MOTA | 837 | 0 | GLY | 56 | -10.473 | 13.543 -15. | 294 1 | 1.00 | 0.42 |
| MOTA | 838 | N | ILE | 57 | -12.613 | 13.078 -15. | _ | | |
| ATOM | 839 | | | | _ | | | 1.00 | 0.37 |
| - | | HN | ILE | 57 | -13.533 | 13.275 -15. | 380 1 | 1.00 | 0.39 |
| MOTA | 840 | CA | ILE | 57 | -12.352 | 12.002 -14. | | 1.00 | |
| MOTA | 841 | HA | ILE | | | | | | 0.35 |
| | | | | 57 | -11.406 | 12.184 -13. | 616 | 1.00 | 0.38 |
| ATOM | 842 | CB | ILE | 57 | -13.473 | 12.000 -13. | 064 | 1.00 | 0.41 |
| ATOM | 843 | HB | ILE | 57 | -14.415 | | | | |
| | | | | | | 11.820 -13. | | 1.00 | 0.42 |
| ATOM | 844 | CG1 | ILE | .57 | -13.508 | 13.363 -12. | 360 1 | 1.00 | 0.48 |
| MOTA | 845 | HG11 | ILE | 57 | -13.512 | 14.148 -13. | | | |
| ATOM | 846 | | | | | | | 1.00 | 0.48 |
| | _ | HG12 | ILE | 57 | -12.631 | 13.465 -11. | 737 1 | 1.00 | 0.51 |
| ATOM | 847 | CG2 | ILE | 57 | -13.216 | 10.896 -12. | | 1.00 | |
| ATOM | 848 | HG21 | | | | | | | 0.44 |
| | | | | 57 | -13.315 | 9.932 -12. | 513 | 1.00 | 1.19 |
| MOTA | 849 | HG22 | ILE | 57 | -13.934 | 10.977 -11. | 235 1 | 1.00 | 1.09 |
| ATOM | 850 | HG23 | ILE | 57 | | | - | | _ |
| | | | | | -12.218 | 11.000 -11. | | 1.00 | 1.04 |
| MOTA | 851 | CD1 | ILE | 57 | -14.765 | 13.484 -11. | 488 1 | 1.00 | 0.56 |
| MOTA | 852 | HD11 | ILE | 57 | -15.459 | 12.693 -11. | - | | |
| ATOM | | | | | | | | 1.00 | 1.08 |
| | 853 | HD12 | ILE | 57 | -15.235 | 14.439 -11. | 668 1 | 1.00 | 1.24 |
| ATOM | 854 | HD13 | ILE | · 57 | -14.487 | 13.413 -10. | | 1.00 | |
| ATOM | 855 | C | | | | | | | 1.14 |
| | | _ | ILE | 57 | -12.307 | 10.647 - 14. | 817 | 1.00 | 0.30 |
| MOTA | 856 | 0 | ILE | 57 | -13.139 | 10.353 -15. | 653 1 | 1.00 | 0.31 |
| ATOM | 857 | N | ALA | 58 | | _ | : : : : · | | |
| _ | | | | | -11.337 | 9.828 -14. | | 1.00 | 0.26 |
| ATOM | 858 | HN | ALA | 58 | -10.679 | 10.096 -13. | 817 | 1.00 | 0.27 |
| ATOM | 859 | CA | ALA | 58 | -11.221 | 8.489 -15. | | | |
| ATOM | 860 | HA | | | | | | 1.00 | 0.23 |
| | | | ALA | 58 | -11.957 | 8.398 -15. | 932 | 1.00 | 0.25 |
| MOTA | 861 | CB | ALA | 58 | -9.824 | 8.339 -15. | 749 1 | 1.00 | 0.23 |
| ATOM | 862 | HB1 | ALA | 58 | -9.843 | _ | | | |
| MOTA | | _ | | | | 7.585 -16. | | 1.00 | 0.97 |
| | 863 | | ALA | 58 | -9.129 | 8.044 -14. | 976 3 | 1.00 | 1.11 |
| MOTA | 864 | HB3 | ALA | 58 | -9.513 | 9.280 -16. | | 1.00 | |
| ATOM | 865 | C | ALA | 58 | | | | | 1.03 |
| | | | | | -11.443 | 7.387 -14. | _ | 1.00 | 0.23 |
| MOTA | 866 | 0 | ALA | 58 | -11.389 | 7.617 -12. | | 1.00 | 0.27 |
| MOTA | 867 | N | ASP | 59 | -11.701 | 6.189 -14. | | 1.00 | |
| ATOM | 868 | | | | | | | | 0.25 |
| | | HN | ASP | 59 | -11.744 | 6.028 -15. | 530 1 | 1.00 | 0.28 |
| MOTA | 869 | CA | ASP | 59 | -11.934 | 5.069 -13. | | | |
| ATOM | 870 | HA | ASP | | | | | 1.00 | 0.27 |
| | | | | 59 | -12.788 | 5.296 -12. | | 1.00 | 0.34 |
| MOTA | 871 | CB | ASP | 59 | -12.207 | 3.785 -14. | | 1.00 | 0.33 |
| MOTA | 872 | | | 59 | | | | | |
| | | | | | -12.203 | 2.942 - 13. | | 1.00 | 0.34 |
| MOTA | 873 | | ASP | 59 | -11.438 | 3.651 -15. | | 1.00 | 0.32 |
| MOTA | 874 | CG | ASP | 59 | -13.572 | | | | |
| ATOM | | | | | | 3.880 -15. | | 1.00 | 0.44 |
| | 875 | | ASP | 59 | -13.791 | 3.139 -16. | 028 | 1.00 | 1.20 |
| MOTA | 876 | OD2 | ASP | 59 | -14.374 | 4.691 -14. | | | |
| MOTA | 877 | C | | | | | | 1.00 | 1.14 |
| | | | ASP | 59 | -10.700 | 4.863 -12. | | 1.00 | 0.22 |
| MOTA | 878 | 0 | ASP | 59 | -10.806 | 4.767 -11. | | 1.00 | 0.27 |
| MOTA | 879 | N | ILE | 60 | | | | | |
| | | | | | -9.534 | 4.780 -13. | | 1.00 | 0.18 |
| MOTA | 880 | HN | ILE | 60 | -9.478 | 4.850 -14. | | 1.00 | 0.20 |
| ATOM | 881 | CA | ILE | 60 | -8.291 | 4.561 -12. | | | |
| MOTA | 882 | | | | | | | 1.00 | 0.22 |
| | | HA | ILE | 60 | -8.554 | 4.303 -11. | 512 | 1.00 | 0.28 |
| MOTA | 883 | CB | ILE | 60 | -7.502 | 3.404 -13. | | 1.00 | |
| ATOM | 884 | HB | ILE | 60 | | | | | 0.27 |
| | | | | | -7.255 | 3.655 -14. | | 1.00 | 0.28 |
| ATOM | 885 | CG1 | ILE | 60 | -8.377 | 2.146 -13. | | 1.00 | 0.30 |
| - | | | | | - | | | | • ••• |

| ATOM | 887 | HG12 | ILE | 60 | -8.541 | 1.839 | -12.113 | 1.00 | 0.36 |
|------|-----|------|-----|------|---------------|--------|--------------|------|------|
| MOTA | 888 | CG2 | | 60 | -6.210 | | -12.369 | 1.00 | 0.39 |
| ATOM | | | ILE | | | | | | |
| | | | | 60 | -6.456 | _ | -11.409 | 1.00 | 1.05 |
| MOTA | | | | 60 | -5.658 | 4.043 | -12.228 | 1.00 | 1.10 |
| MOTA | 891 | HG23 | ILE | 60 | -5.600 | 2.428 | -12.921 | 1.00 | 1.12 |
| MOTA | 892 | CD1 | ILE | 60 | -7.688 | 1.015 | -13.904 | 1.00 | 0.38 |
| MOTA | _ | | | 60 | -7.209 | | -14.786 | 1.00 | 1.07 |
| MOTA | | | ILE | 60 | | | | | |
| | | | | | -8.424 | | -14.196 | 1.00 | 1.14 |
| MOTA | | HD13 | ILE | 60 | -6.948 | | -13.270 | 1.00 | 1.04 |
| MOTA | 896 | Ç | ILE | 60 | -7.438 | 5.834 | -12.518 | 1.00 | 0.20 |
| ATOM | 897 | 0 | ILE | 60 | -6.731 | 6.115 | -13.464 | 1.00 | 0.25 |
| MOTA | 898 | N | MET | 61 | -7.473 | | -11.448 | 1.00 | 0.20 |
| ATOM | 899 | HN | MET | 61 | -8.033 | | -10.687 | 1.00 | |
| | 900 | | | | | | | | 0.25 |
| MOTA | | CA | MET | 61 | -6.641 | | -11.373 | 1.00 | 0.20 |
| MOTA | 901 | | MET | 61 | -6.327 | | -12.366 | 1.00 | 0.19 |
| MOTA | 902 | CB | MET | 61 | -7.464 | 8.963 | -10.773 | 1.00 | 0.24 |
| MOTA | 903 | HB1 | MET | 61 | -8.331 | 9.137 | -11.392 | 1.00 | 0.35 |
| MOTA | 904 | HB2 | MET | 61 | -6.860 | | -10.743 | 1.00 | 0.33 |
| MOTA | 905 | CG | MET | 61 | -7.918 | 8.604 | -9.358 | 1.00 | 0.31 |
| ATOM | 906 | | MET | 61 | -7.146 | 8.870 | | | |
| | 907 | | | | . — | | -8.653 | 1.00 | 0.66 |
| ATOM | | | MET | 61 | -8.112 | 7.544 | -9.300 | 1.00 | 0.67 |
| MOTA | 908 | SD | MET | 61 | -9.433 | 9.519 | -8.967 | 1.00 | 0.54 |
| MOTA | 909 | CE | MET | 61 | -8.878 | 11.154 | -9.516 | 1.00 | 0.40 |
| MOTA | 910 | HE1 | MET | 61 | -9.492 | 11.914 | -9.056 | 1.00 | 1.06 |
| MOTA | 911 | HE2 | MET | 61 | -8.968 | 11.227 | -10.589 | 1.00 | 1.16 |
| ATOM | 912 | HE3 | MET | 61 | -7.846 | 11.298 | -9.232 | 1.00 | |
| ATOM | 913 | C | MET | 61 | | | | | 1.12 |
| | | | | | -5.396 | | -10.524 | 1.00 | 0.20 |
| ATOM | 914 | 0 | MET | 61 | -5.478 | 6.951 | -9.463 | 1.00 | 0.22 |
| MOTA | 915 | N | ILE | 62 | -4.241 | 7.937 | -11.001 | 1.00 | 0.20 |
| ATOM | 916 | HN | ILE | 62 | -4.207 | 8.393 | -11.868 | 1.00 | 0.21 |
| ATOM | 917 | CA | ILE | 62 | -2.971 | 7.678 | -10.252 | 1.00 | 0.21 |
| MOTA | 918 | HA | ILE | 62 | -3.156 | 6.982 | | 1.00 | 0.20 |
| ATOM | 919 | CB | ILE | | | | | _ | |
| | | | | | -1.938 | | -11.211 | 1.00 | 0.24 |
| MOTA | 920 | HB | ILE | 62 | -1.753 | | -12.012 | 1.00 | 0.26 |
| MOTA | 921 | CG1 | ILE | 62 | -2.480 | 5.762 | -11.785 | 1.00 | 0.23 |
| MOTA | 922 | HG11 | ILE | 62 | -3.479 | 5.922 | -12.162 | 1.00 | 0.20 |
| MOTA | 923 | HG12 | ILE | 62 | -2.508 | 5.018 | -11.003 | 1.00 | 0.24 |
| MOTA | 924 | CG2 | ILE | 62 | -0.635 | | -10.455 | 1.00 | 0.30 |
| ATOM | 925 | HG21 | | 62 | -0.863 | | -9.466 | 1.00 | |
| ATOM | 926 | HG22 | | 62 | | | | | 1.08 |
| | | | | | -0.070 | | -10.375 | 1.00 | 1.12 |
| MOTA | 927 | HG23 | ILE | 62 | -0.052 | | -10.988 | 1.00 | 0.99 |
| ATOM | 928 | CD1 | | 62 | -1.584 | 5.262 | -12.927 | 1.00 | 0.29 |
| MOTA | 929 | HD11 | ILE | 62 | -0.979 | 6.073 | -13.305 | 1.00 | 1.02 |
| MOTA | 930 | HD12 | ILE | 62 | -2.201 | | -13.724 | 1.00 | 1.09 |
| MOTA | 931 | HD13 | ILE | 62 | -0.941 | | -12.559 | 1.00 | 1.07 |
| ATOM | 932 | C | ILE | 62 | -2.423 | 8.988 | -9.677 | _ | |
| MOTA | 933 | ŏ | ILE | 62 | | | | 1.00 | 0.22 |
| MOTA | | _ | | | -2.393 | | -10.343 | 1.00 | 0.27 |
| | 934 | N | SER | 63 | -1.993 | 8.976 | | 1.00 | 0.20 |
| ATOM | 935 | HN | SER | 63 | -2.028 | 8.147 | | 1.00 | 0.18 |
| MOTA | 936 | CA | SER | 63 | -1.452 | 10.226 | -7.829 | 1.00 | 0.22 |
| MOTA | 937 | HA | SER | 63 | -0.998 | 10.836 | -8.597 | 1.00 | 0.26 |
| MOTA | 938 | CB | SER | 63 | -2.597 | 11.000 | -7.176 | 1.00 | 0.24 |
| MOTA | 939 | HB1 | SER | 63 | -3.448 | 11.012 | -7.845 | 1.00 | 0.25 |
| MOTA | 940 | HB2 | SER | 63 | -2.286 | | | _ | |
| MOTA | 941 | OG | | | | 12.012 | -6.978 | 1.00 | 0.29 |
| ATOM | | | SER | 63 | -2.951 | 10.369 | | 1.00 | 0.25 |
| | 942 | HG | SER | 63 | -3.682 | 9.772 | -6.127 | 1.00 | 0.85 |
| ATOM | 943 | C | SER | 63 | -0.404 | 9.879 | -6.764 | 1.00 | 0.21 |
| MOTA | 944 | 0 | SER | 63 | -0.364 | 8.775 | -6.259 | 1.00 | 0.20 |
| ATOM | 945 | N | PHE | 64 | 0.440 | 10.823 | | 1.00 | 0.24 |
| ATOM | 946 | HN | PHE | 64 | 0.380 | 11.705 | _ | 1.00 | 0.27 |
| ATOM | 947 | CA | PHE | 64 | 1.490 | 10.569 | | 1.00 | 0.24 |
| ATOM | 948 | HA | PHE | 64 | | | | | |
| ATOM | 949 | | | | 1.560 | 9.511 | - | 1.00 | 0.22 |
| | | CB | PHE | 64 | 2.840 | 11.084 | | 1.00 | 0.28 |
| ATOM | 950 | HB1 | _ | 64 | 3.564 | 11.047 | | 1.00 | 0.32 |
| MOTA | 951 | HB2 | | 64 | 2.730 | 12.103 | -6.235 | 1.00 | 0.32 |
| MOTA | 952 | CG | PHE | 64 | 3.316 | 10.220 | | 1.00 | 0.28 |
| MOTA | 953 | CD1 | PHE | - 64 | 4.112 | 9.096 | _ | 1.00 | 0.30 |
| ATOM | 954 | HD1 | PHE | 64 | 4.385 | 8.844 | -5.774 | 1.00 | 0.32 |
| MOTA | 955 | | PHE | 64 | 2.963 | | | | |
| ATOM | 956 | | | | | 10.545 | | 1.00 | 0.33 |
| | | | PHE | 64 | 2.350 | 11.412 | | 1.00 | 0.37 |
| ATOM | 957 | CE1 | PHE | 64 | 4.553 | 8.297 | | 1.00 | 0.36 |
| ATOM | 958 | HE1 | | 64 | 5.166 | 7.430 | | 1.00 | 0.40 |
| MOTA | 959 | CE2 | | 64 | 3.403 | 9.747 | -9.417 | 1.00 | 0.40 |
| MOTA | 960 | HE2 | PHE | 64 | 3.130 | 9.998 | | 1.00 | 0.47 |
| ATOM | 961 | CZ | PHE | 64 | 4.198 | 8.623 | -9.165 | 1.00 | 0.40 |
| ATOM | 962 | HZ | PHE | 64 | 4.538 | 8.007 | | 1.00 | 0.47 |
| MOTA | 963 | C | PHE | 64 | 1.115 | 11.318 | -4.097 | 1.00 | 0.47 |
| | | _ | | | | ~~.710 | ~6.047 | | 0.21 |

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| | | _ | | | _ | | | | |
|---------|------|------|-----|----------|--------|--------|--------|------|------|
| ATOM | 964 | | PHE | 64 | 0.924 | 12.518 | -4.108 | 1.00 | 0.36 |
| MOTA | 965 | N | GLY | - 65 | 0.996 | 10.617 | -2.996 | 1.00 | 0.30 |
| MOTA | 966 | HN | GLY | 65 | 1.146 | 9.649 | -3.017 | 1.00 | 0.33 |
| ATOM | 967 | CA | GLY | 65 | 0.615 | 11.282 | -1.709 | 1.00 | 0.38 |
| ATOM | 968 | | GLY | 65 | -0.152 | 10.697 | -1.224 | 1.00 | 0.46 |
| MOTA | 969 | | GLY | 65 | 0.230 | 12.270 | -1.913 | | |
| ATOM | 970 | | GLY | 65 | | | | 1.00 | 0.45 |
| | 971 | | | | 1.823 | 11.397 | -0.770 | 1.00 | 0.32 |
| MOTA | | | GLY | 65 | 2.926 | 11.007 | -1.098 | 1.00 | 0.40 |
| MOTA | 972 | N | ILE | 66 | 1.598 | 11.926 | 0.408 | 1.00 | 0.30 |
| MOTA | 973 | HN | ILE | 66 | 0.691 | 12.220 | 0.635 | 1.00 | 0.36 |
| ATOM | 974 | CA | ILE | 66 | 2.691 | 12.081 | 1.417 | 1.00 | 0.36 |
| MOTA | 975 | HA | ILE | 66 | 3.564 | 11.534 | 1.093 | 1.00 | 0.40 |
| MOTA | 976 | CB | ILE | 66 | 3.040 | 13.564 | 1.571 | 1.00 | 0.41 |
| ATOM | 977 | HB | ILE | 66 | 2.127 | 14.134 | 1.656 | 1.00 | 0.64 |
| MOTA | 978 | CG1 | ILE | 66 | 3.829 | 14.026 | 0.337 | | |
| ATOM | | HG11 | ILE | 66 | | | | 1.00 | 0.68 |
| MOTA | | | | | 3.301 | 13.729 | -0.557 | 1.00 | 0.95 |
| | | HG12 | ILE | 66 | 4.804 | 13.561 | 0.346 | 1.00 | 1.01 |
| ATOM | 981 | CG2 | ILE | 66 | 3.886 | 13.764 | 2.831 | 1.00 | 0.93 |
| MOTA | | HG21 | ILE | 66 | 4.372 | 14.727 | 2.790 | 1.00 | 1.50 |
| ATOM | 983 | HG22 | ILE | 66 | 4.632 | 12.986 | 2.891 | 1.00 | 1.41 |
| MOTA | 984 | HG23 | ILE | 66 | 3.249 | 13.720 | 3.702 | 1.00 | 1.54 |
| ATOM | 985 | CD1 | ILE | 66 | 3.997 | 15.551 | 0.343 | 1.00 | 0.70 |
| ATOM | 986 | HD11 | ILE | 66 | 4.944 | 15.806 | 0.797 | 1.00 | 1.22 |
| ATOM | 987 | HD12 | ILE | 66 | 3.196 | 16.009 | 0.902 | 1.00 | 1.28 |
| ATOM | 988 | HD13 | ILE | 66 | 3.979 | 15.917 | -0.673 | 1.00 | |
| MOTA | 989 | C | ILE | 66 | | | | | 1.23 |
| MOTA | 990 | | | | 2.207 | 11.519 | 2.760 | 1.00 | 0.46 |
| | | 0 | ILE | 66 | 1.021 | 11.363 | 2.958 | 1.00 | 0.54 |
| MOTA | 991 | N | LYS | 67 | 3.129 | 11.205 | 3.659 | 1.00 | 0.59 |
| MOTA | 992 | HN | LYS | 67 | 4.073 | 11.343 | 3.434 | 1.00 | 0.64 |
| MOTA | 993 | CA | LYS | 67 | 2.780 | 10.630 | 5.014 | 1.00 | 0.74 |
| MOTA | 994 | HA | LYS | 67 | 3:072 | 9.594 | 5.038 | 1.00 | 0.83 |
| MOTA | 995 | CB | LYS | 67 | 3.550 | 11.404 | 6.102 | 1.00 | 0.90 |
| MOTA | 996 | HB1 | LYS | 67 | 3.237 | 12.438 | 6.089 | 1.00 | 0.89 |
| ATOM | 997 | HB2 | LYS | 67 | 4.608 | 11.352 | 5.891 | 1.00 | 0.96 |
| MOTA | 998 | CG | LYS | 67 | 3.287 | 10.815 | 7.504 | 1.00 | 1.08 |
| MOTA | 999 | | LYS | 67 | 2.254 | 10.524 | 7.598 | | |
| MOTA | 1000 | | LYS | 67 | 3.510 | | | 1.00 | 1.31 |
| ATOM | 1001 | CD | LYS | | | 11.565 | 8.249 | 1.00 | 1.33 |
| | | | | 67 | 4.179 | 9.590 | 7.746 | 1.00 | 0.98 |
| ATOM | 1002 | | LYS | 67 | 5.216 | 9.885 | 7.694 | 1.00 | 1.07 |
| MOTA | 1003 | | LYS | 67 | 3.979 | 8.839 | 6.999 | 1.00 | 1.07 |
| MOTA | 1004 | CE | LYS | 67 | 3.885 | 9.016 | 9.135 | 1.00 | 1.17 |
| ATOM | 1005 | | LYS | 67 | 4.331 | 8.036 | 9.220 | 1.00 | 1.64 |
| ATOM | 1006 | HE2 | LYS | 67 | 2.817 | 8.938 | 9.272 | 1.00 | 1.50 |
| ATOM | 1007 | NZ | LYS | 67 | 4.453 | 9.913 | 10.180 | 1.00 | 1.93 |
| ATOM | 1008 | HZ1 | LYS | 67 | 4.569 | 10.870 | 9.792 | 1.00 | 2.38 |
| MOTA | 1009 | HZ2 | LYS | 67 | 5.378 | 9.547 | 10.485 | 1.00 | 2.43 |
| ATOM | 1010 | HZ3 | LYS | 67 | 3.808 | 9.948 | 10.995 | 1.00 | 2.40 |
| MOTA | 1011 | C | LYS | 67 | 1.274 | 10.732 | 5.280 | 1.00 | 0.72 |
| MOTA | 1012 | Ö | LYS | 67 | 0.530 | 9.804 | | | |
| ATOM | 1013 | N | GLU | 68 | 0.815 | | 5.035 | 1.00 | 0.79 |
| ATOM | 1014 | HN | | | | 11.855 | 5.760 | 1.00 | 0.77 |
| | | | GLU | 68 | 1.425 | 12.601 | 5.939 | 1.00 | 0.84 |
| MOTA | 1015 | CA | GLU | 68 | -0.645 | 12.004 | 6.011 | 1.00 | 0.84 |
| ATOM | 1016 | HA | GLU | 68 | -1.014 | 11.130 | 6.530 | 1.00 | 0.99 |
| MOTA | 1017 | CB | GLU | 68 | -0.895 | 13.254 | 6.860 | 1.00 | 1.05 |
| MOTA | 1018 | HB1 | GLU | 68 | -0.393 | 13.149 | 7.810 | 1.00 | 1.23 |
| MOTA | 1019 | HB2 | GLU | 68 | -1.956 | 13.370 | 7.024 | 1.00 | 1.10 |
| ATOM | 1020 | CG | GLU | 68 | -0.353 | 14.487 | 6.134 | 1.00 | 1.15 |
| MOTA | 1021 | HG1 | GLU | 68 | -1.000 | 14.730 | 5.304 | 1.00 | 1.32 |
| MOTA | 1022 | HG2 | GLU | 68 | 0.642 | 14.281 | 5.768 | 1.00 | 1.28 |
| MOTA | 1023 | CD | GLU | 68 | -0.308 | 15.669 | 7.104 | 1.00 | 1.75 |
| MOTA | 1024 | OE1 | | 68 | 0.246 | 16.692 | 6.736 | | |
| MOTA | 1025 | | | 68 | -0.823 | | | 1.00 | 2.45 |
| ATOM | 1026 | | GLU | 68 | | 15.530 | 8.202 | 1.00 | 2.16 |
| MOTA | 1027 | | | | -1.346 | 12.132 | 4.660 | 1.00 | 0.76 |
| | | | GLU | 68 | -0.899 | 12.859 | 3.795 | 1.00 | 1.11 |
| ATOM | 1028 | | HIS | 69 | -2.420 | 11.414 | 4.454 | 1.00 | 0.94 |
| ATOM | 1029 | | HIS | 69 | -2.755 | 10.815 | 5.155 | 1.00 | 1.32 |
| MOTA | 1030 | | HIS | 69 | -3.114 | 11.487 | 3.136 | 1.00 | 1.04 |
| MOTA | 1031 | | HIS | 69 | -2.877 | 12.437 | 2.679 | 1.00 | 1.25 |
| MOTA | 1032 | CB | HIS | 69 | -2.545 | 10.358 | 2.243 | 1.00 | 1.49 |
| ATOM | 1033 | | | 69 | -1.750 | 9.862 | 2.783 | 1.00 | 2.12 |
| MOTA | 1034 | | | 69 | -2.131 | 10.798 | 1.351 | 1.00 | 2.12 |
| MOTA | 1035 | | HIS | 69 | -3.570 | 9.333 | 1.837 | | |
| MOTA | 1036 | | | 69 | -3.818 | | | 1.00 | 0.95 |
| ATOM | 1037 | | HIS | 69 | | 8.195 | 2.588 | 1.00 | 1.43 |
| MOTA | 1038 | | HIS | | -3.415 | 7.972 | 3.453 | 1.00 | 1.83 |
| ATOM | 1039 | | HIS | 69 69 | -4.355 | 9.223 | 0.717 | 1.00 | 1.04 |
| MOTA | 1040 | | | 69 60 | -4.403 | 9.946 | -0.082 | 1.00 | 1.41 |
| -10 Vii | 7040 | CEI | HIS | 69 | -4.715 | 7.452 | 1.912 | 1.00 | 1.81 |

| > ==== | 1041 | | | | | | | | _ |
|--------|------|-----|-----|----|---------|--------|--------|------|------|
| ATOM | 1041 | | HIS | 69 | -5.097 | 6.502 | 2.257 | 1.00 | 2.54 |
| MOTA | 1042 | NE2 | HIS | 69 | -5.075 | 8.032 | 0.765 | 1.00 | 1.53 |
| MOTA | 1043 | C | HIS | 69 | -4.643 | 11.435 | 3.341 | 1.00 | 1.14 |
| MOTA | 1044 | 0 | HIS | 69 | -5.392 | 10.889 | 2.556 | 1.00 | 1.76 |
| MOTA | 1045 | N | GLY | 70 | -5.108 | 12.065 | 4.393 | 1.00 | 1.49 |
| MOTA | 1046 | HN | GLY | 70 | -4.487 | 12.532 | 4.990 | 1.00 | 1.98 |
| MOTA | 1047 | CA | GLY | 70 | -6.576 | 12.123 | 4.665 | 1.00 | 1.86 |
| MOTA | 1048 | HA1 | GLY | 70 | -7.071 | 12.633 | 3.852 | 1.00 | 2.28 |
| MOTA | 1049 | HA2 | GLY | 70 | -6.746 | 12.667 | | | |
| MOTA | 1050 | C | GLY | 70 | | | 5.583 | 1.00 | 2.09 |
| | | _ | | | -7.155 | 10.716 | 4.801 | 1.00 | 1.81 |
| ATOM | 1051 | 0 | GLY | 70 | -8.182 | 10.404 | 4.232 | 1.00 | 2.53 |
| MOTA | 1052 | N | ASP | 71 | -6.513 | 9.863 | 5.545 | 1.00 | 1.55 |
| MOTA | 1053 | HN | ASP | 71 | -5.686 | 10.127 | 5.999 | 1.00 | 1.66 |
| MOTA | 1054 | CA | ASP | 71 | -7.047 | 8.484 | 5.701 | 1.00 | 1.91 |
| MOTA | 1055 | AH | ASP | 71 | -8.126 | 8.513 | 5.684 | 1.00 | 2.42 |
| ATOM | 1056 | CB | ASP | 71 | -6.546 | 7.620 | 4.546 | 1.00 | 2.67 |
| MOTA | 1057 | HB1 | ASP | 71 | -6.623 | 6.578 | 4.813 | 1.00 | 3.03 |
| MOTA | 1058 | HB2 | ASP | 71 | -5.514 | 7.865 | 4.341 | 1.00 | 2.88 |
| ATOM | 1059 | CG | ASP | 71 | -7.397 | 7.892 | 3.303 | 1.00 | 3.56 |
| MOTA | 1060 | OD1 | | 71 | -8.476 | 7.330 | 3.215 | 1.00 | 4.08 |
| MOTA | 1061 | OD2 | ASP | 71 | -6.960 | 8.664 | 2.465 | 1.00 | |
| ATOM | 1062 | C | ASP | 71 | -6.577 | | | | 4.16 |
| ATOM | 1063 | ŏ | ASP | 71 | | 7.889 | 7.028 | 1.00 | 1.46 |
| ATOM | 1064 | N | | | -5.600 | 8.323 | 7.605 | 1.00 | 1.78 |
| | | | PHE | 72 | -7.260 | 6.886 | 7.507 | 1.00 | 1.36 |
| MOTA | 1065 | HN | PHE | 72 | -8.038 | 6.546 | 7.018 | 1.00 | 1.67 |
| ATOM | 1066 | CA | PHE | 72 | -6.849 | 6.248 | 8.786 | 1.00 | 1.48 |
| MOTA | 1067 | HA | PHE | 72 | -6.504 | 7.007 | 9.473 | 1.00 | 1.75 |
| MOTA | 1068 | CB | PHE | 72 | -8.037 | 5.503 | 9.399 | 1.00 | 2.01 |
| MOTA | 1069 | HB1 | PHE | 72 | -8.374 | 6.028 | 10.281 | 1.00 | 2.58 |
| MOTA | 1070 | HB2 | PHE | 72 | -7.733 | 4.503 | 9.669 | 1.00 | 2.43 |
| ATOM | 1071 | CG | PHE | 72 | -9.161 | 5.434 | 8.395 | 1.00 | 2.30 |
| ATOM | 1072 | CD1 | | 72 | -9.414 | 4.243 | 7.704 | 1.00 | |
| | 1073 | HD1 | | 72 | -8.802 | | | | 2.86 |
| ATOM | 1074 | | PHE | 72 | | 3.372 | 7.887 | 1.00 | 3.09 |
| ATOM | | | | | -9.954 | 6.563 | 8.158 | 1.00 | 2.97 |
| | 1075 | HD2 | | 72 | -9.758 | 7.482 | 8.691 | 1.00 | 3.28 |
| MOTA | 1076 | CE1 | PHE | 72 | -10.459 | 4.182 | 6.775 | 1.00 | 3.73 |
| ATOM | 1077 | HE1 | | 72 | -10.655 | 3.264 | 6.242 | 1.00 | 4.46 |
| MOTA | 1078 | CE2 | PHE | 72 | -10.999 | 6.502 | 7.229 | 1.00 | 3.80 |
| MOTA | 1079 | HE2 | PHE | 72 | -11.610 | 7.374 | 7.045 | 1.00 | 4.54 |
| MOTA | 1080 | CZ | PHE | 72 | -11.252 | 5.312 | 6.537 | 1.00 | 4.08 |
| ATOM | 1081 | HZ | PHE | 72 | -12.058 | 5.264 | 5.821 | 1.00 | 4.92 |
| MOTA | 1082 | C | PHE | 72 | -5.716 | 5.266 | 8.500 | 1.00 | |
| MOTA | 1083 | ŏ | PHE | 72 | -5.384 | | | | 1.41 |
| ATOM | 1084 | N | TYR | 73 | | 4.430 | 9.318 | 1.00 | 2.20 |
| ATOM | 1085 | HN | | | -5.120 | 5.371 | 7.338 | 1.00 | 1.12 |
| ATOM | | | TYR | 73 | -5.412 | 6.059 | 6.703 | 1.00 | 1.48 |
| | 1086 | CA | TYR | 73 | -3.999 | 4.457 | 6.972 | 1.00 | 1.25 |
| MOTA | 1087 | HA | TYR | 73 | -3.774 | 3.793 | 7.790 | 1.00 | 1.46 |
| ATOM | 1088 | CB | TYR | 73 | -4.391 | 3.635 | 5.742 | 1.00 | 1.86 |
| ATOM | 1089 | HB1 | | 73 | -3.531 | 3.082 | 5.395 | 1.00 | 2.35 |
| MOTA | 1090 | HB2 | TYR | 73 | -4.726 | 4.300 | 4.961 | 1.00 | 2.46 |
| MOTA | 1091 | CG | TYR | 73 | -5.498 | 2.670 | 6.089 | 1.00 | 2.08 |
| MOTA | 1092 | CD1 | TYR | 73 | -5.241 | 1.585 | 6.934 | 1.00 | 2.58 |
| MOTA | 1093 | HD1 | TYR | 73 | -4.252 | 1.444 | 7.347 | 1.00 | 2.82 |
| MOTA | 1094 | CD2 | TYR | 73 | -6.779 | 2.853 | 5.553 | 1.00 | 2.85 |
| MOTA | 1095 | HD2 | | 73 | -6.978 | 3.691 | 4.901 | 1.00 | 3.24 |
| MOTA | 1096 | CE1 | | 73 | -6.264 | 0.683 | 7.244 | 1.00 | |
| ATOM | 1097 | HE1 | | 73 | -6.066 | -0.155 | 7.896 | _ | 3.48 |
| MOTA | 1098 | CE2 | | 73 | -7.802 | 1.952 | | 1.00 | 4.19 |
| MOTA | 1099 | HE2 | | 73 | | | 5.865 | 1.00 | 3.68 |
| ATOM | 1100 | | | | -8.789 | 2.093 | 5.452 | 1.00 | 4.49 |
| | | CZ | TYR | 73 | -7.545 | 0.866 | 6.710 | 1.00 | 3.90 |
| MOTA | 1101 | OH | TYR | 73 | -8.554 | -0.024 | 7.013 | 1.00 | 5.00 |
| MOTA | 1102 | HH | TYR | 73 | -8.689 | -0.590 | 6.249 | 1.00 | 5.22 |
| ATOM | 1103 | C | TYR | 73 | -2.755 | 5.273 | 6.609 | 1.00 | 0.95 |
| MOTA | 1104 | 0 | TYR | 73 | -2.219 | 5.127 | 5.529 | 1.00 | 1.21 |
| MOTA | 1105 | N | PRO | 74 | -2.273 | 6.106 | 7.495 | 1.00 | 0.74 |
| MOTA | 1106 | CA | PRO | 74 | -1.054 | 6.895 | 7.197 | 1.00 | 0.82 |
| MOTA | 1107 | HA | PRO | 74 | -1.254 | 7.648 | 6.453 | 1.00 | 1.05 |
| MOTA | 1108 | CB | PRO | 74 | -0.746 | 7.558 | 8.543 | | |
| ATOM | 1109 | HB1 | | 74 | -0.786 | 8.631 | | 1.00 | 1.18 |
| MOTA | 1110 | HB2 | | 74 | 0.239 | | 8.438 | 1.00 | 1.46 |
| ATOM | 1111 | CG | PRO | 74 | | 7.261 | 8.876 | 1.00 | 1.28 |
| ATOM | 1112 | | | | -1.795 | 7.105 | 9.566 | 1.00 | 1.35 |
| | | HG1 | | 74 | -2.229 | 7.967 | 10.049 | 1.00 | 1.70 |
| MOTA | 1113 | HG2 | | 74 | -1.330 | 6.468 | 10.305 | 1.00 | 1.61 |
| ATOM | 1114 | CD | PRO | 74 | -2.889 | 6.328 | 8.828 | 1.00 | 1.04 |
| ATOM | 1115 | HD2 | | 74 | -3.098 | 5.393 | 9.328 | 1.00 | 1.24 |
| MOTA | 1116 | HD1 | | 74 | -3.778 | 6.929 | 8.733 | 1.00 | 1.14 |
| MOTA | 1117 | C | PRO | 74 | 0.097 | 5.988 | 6.765 | 1.00 | 0.65 |
| | | | | | | | | | |

| MOTA | 1118 | 0 | PRO | 74 | 0 176 | 4 922 | 7 106 | 1 00 | 0 55 |
|--------------|--------------|----------|------------|------------|----------------|----------------|----------------|------|--------------|
| MOTA | 1119 | N | PHE | 7 5 | 0.136 | 4.822 | 7.106 | 1.00 | 0.66 0.56 |
| MOTA | 1120 | HN | PHE | 75 75 | 1,038 1.000 | 6.503 7.447 | 6.032 5.770 | 1.00 | 0.61 |
| ATOM | 1121 | CA | PHE | 75 75 | 2.179 | 5.651 | 5.605 | 1.00 | 0.45 |
| ATOM | 1122 | HA | PHE | · 75 | 1.816 | 4.659 | 5.360 | 1.00 | 0.48 |
| ATOM | 1123 | СВ | PHE | 75 75 | 2.859 | 6.266 | 4.379 | 1.00 | 0.42 |
| MOTA | 1124 | HB1 | PHE | ,5 75 | 3.761 | 5.718 | 4.153 | 1.00 | 0.44 |
| MOTA | 1125 | HB2 | PHE | .5 75 | 3.104 | 7.298 | 4.582 | 1.00 | 0.45 |
| MOTA | 1126 | CG | PHE | 75 | 1.915 | 6.190 | 3.200 | 1.00 | 0.48 |
| MOTA | 1127 | CD1 | PHE | 75 | 1.764 | 4.986 | 2.501 | 1.00 | 0.41 |
| ATOM | 1128 | HD1 | PHE | 75 | 2.329 | 4.115 | 2.797 | 1.00 | 0.45 |
| MOTA | 1129 | CD2 | PHE | 75 | 1.184 | 7.320 | 2.812 | 1.00 | 0.74 |
| MOTA | 1130 | HD2 | PHE | 75 | 1.300 | 8.249 | 3.349 | 1.00 | 0.90 |
| ATOM | 1131 | CE1 | PHE | 75 | 0.882 | 4.911 | 1.415 | 1.00 | 0.50 |
| MOTA | 1132 | HE1 | PHE | 75 | 0.767 | 3.982 | 0.877 | 1.00 | 0.53 |
| ATOM | 1133 | CE2 | PHE | 75 | 0.304 | 7.245 | 1.724 | 1.00 | 0.85 |
| MOTA | 1134 | HE2 | PHE | 75 | -0.258 | 8.117 | 1.423 | 1.00 | 1.09 |
| MOTA | 1135 | CZ | PHE | 75 | 0.154 | 6.041 | 1.026 | 1.00 | 0.69 |
| · ATOM | 1136 | HZ | PHE | 75 | -0.526 | 5.983 | 0.188 | 1.00 | 0.80 |
| ATOM | 1137 | C | PHE | 75 | 3.159 | 5.561 | 6.776 | 1.00 | 0.43 |
| ATOM | 1138 | 0 | PHE | 75 | 3.111 | 6.360 | 7.690 | 1.00 | 0.50 |
| ATOM | 1139 | N | ASP | 76 | 4.020 | 4.582 | 6.782 | 1.00 | 0.37 |
| ATOM | 1140 | HN | ASP | 76 | 4.028 | 3.929 | 6.050 | 1.00 | 0.32 |
| ATOM | 1141 | CA | ASP | 76 | 4.967 | 4.432 | 7.927 | 1.00 | 0.43 |
| ATOM | 1142 | HA | ASP | 76 | 4.551 | 4.906 | 8.804 | 1.00 | 0.50 |
| ATOM | 1143 | CB | ASP | 76 | 5.180 | 2.946 | 8.215 | 1.00 | 0.46 |
| MOTA | 1144 | HB1 | ASP | 76 | 4.224 | 2.467 | 8.365 | 1.00 | 0.49 |
| MOTA | 1145 | | ASP | 76 | 5.784 | 2.834 | 9.104 | 1.00 | 0.54 |
| MOTA | 1146 | | ASP | 76 76 | 5.892 | 2.295 | 7.028 | 1.00 | 0.38 |
| MOTA | 1147 | | ASP | 76 26 | 6.468 | 1.236 | 7.218 | 1.00 | 0.45 |
| MOTA | 1148 | OD2 | | 76 | 5.846 | 2.864 | 5.950 | 1.00 | 0.30 |
| MOTA MOTA | 1149 1150 | C | ASP | 76 76 | 6.314 | 5.074 | 7.596 | 1.00 | 0.42 |
| MOTA | 1151 | O N | ASP GLY | 76 77 | 7.314 | 4.770 | 8.216 | 1.00 | 0.54 |
| MOTA | 1152 | HN | GLY | 77 | 6.347 | 5.958 | 6.632 | 1.00 | 0.35 |
| MOTA | 1153 | CA | GLY | 77 | 5.525 7.634 | 6.187 | 6.151 | 1.00 | 0.36 |
| ATOM | 1154 | HA1 | | 77 | 8.378 | 6.625 6.388 | 6.267 | 1.00 | 0.38 |
| MOTA | 1155 | HA2 | | לל | 7.484 | 7.696 | 7.004 6.238 | 1.00 | 0.45 |
| ATOM | 1156 | C | GLY | 77 | 8.084 | 6.131 | 4.884 | 1.00 | 0.44 0.31 |
| ATOM | 1157 | ō | GLY | 77 | 7.262 | 5.767 | 4.068 | 1.00 | 0.37 |
| ATOM | 1158 | Ň | PRO | 78 | 9.370 | 6.117 | 4.603 | 1.00 | 0.33 |
| ATOM | 1159 | CA | PRO | 78 | 9.856 | 5.651 | 3.274 | 1.00 | 0.36 |
| MOTA | 1160 | HA | PRO | 78 | 9.435 | 6.254 | 2.488 | 1.00 | 0.42 |
| ATOM | 1161 | CB | PRO | 78 | 11.364 | 5.903 | 3.359 | 1.00 | 0.46 |
| ATOM | 1162 | HB1 | | 78 | 11.671 | 6.542 | 2.545 | 1.00 | 0.56 |
| MOTA | 1163 | HB2 | PRO | 78 | 11.892 | 4:962 | 3.303 | 1.00 | 0.48 |
| ATOM | 1164 | CG | PRO | 78 | 11.675 | 6.592 | 4.694 | 1.00 | 0.64 |
| MOTA | 1165 | HG1 | PRO | 78 | 11.965 | 7.616 | 4.516 | 1.00 | 0.87 |
| MOTA | 1166 | HG2 | PRO | 78 | 12.478 | 6.068 | 5.194 | 1.00 | 0.83 |
| MOTA | 1167 | CD | PRO | 78 | 10.418 | 6.562 | 5.563 | 1.00 | 0.45 |
| MOTA | 1168 | HD2 | | 78 | 10.535 | 5.848 | 6.369 | 1.00 | 0.48 |
| MOTA | 1169 | HD1 | | 78 | 10.187 | 7.544 | 5.944 | 1.00 | 0.49 |
| MOTA | 1170 | C | PRO | 78 | 9.564 | 4.165 | 3.027 | 1.00 | 0.30 |
| MOTA | 1171 | 0 | PRO | 78 | 8.860 | 3.808 | 2.105 | 1.00 | 0.28 |
| MOTA | 1172 | N | SER | 79 70 | 10.102 | 3.297 | 3.840 | 1.00 | 0.31 |
| MOTA MOTA | 1173 | HN | SER | 79 70 | 10.670 | 3.604 | 4.577 | 1.00 | 0.35 |
| MOTA | 1174 1175 | CA HA | SER SER | 79 79 | 9.855 | 1.837 | 3.647 | 1.00 | 0.30 |
| ATOM | 1176 | CB | SER | 79 79 | 9.916 | 1.599 | 2.595 | 1.00 | 0.30 |
| MOTA | 1177 | HB1 | | 79 | 10.911 | 1.037 | 4.410 | 1.00 | 0.37 |
| MOTA | 1178 | HB2 | | 79 79 | 11.888 | 1.465 | 4.225 | 1.00 | 0.42 |
| ATOM | 1179 | OG | SER | 79 | 10.617 | 0.013 1.080 | 4.076 5.800 | 1.00 | 0.39 |
| ATOM | 1180 | HG | SER | 79 | 11.173 | 1,752 | 6.201 | 1.00 | 0.38 |
| ATOM | 1181 | C | SER | 79 | 8.463 | 1,732 | 4.173 | 1.00 | 0.98 0.27 |
| MOTA | 1182 | ŏ | SER | 79 | 7.888 | 2.183 | 4.971 | 1.00 | 0.25 |
| ATOM | 1183 | N | GLY | 80 | 7.927 | 0.356 | 3.734 | 1.00 | 0.31 |
| ATOM | 1184 | HN | GLY | 80 | 8.420 | -0.200 | 3.734 | 1.00 | 0.37 |
| ATOM | 1185 | CA | GLY | 80 | 6.576 | -0.081 | 4.207 | 1.00 | 0.30 |
| ATOM | 1186 | HA1 | | 80 | 6.224 | 0.586 | 4.977 | 1.00 | 0.31 |
| ATOM | 1187 | HA2 | | 80 | 6.646 | -1.083 | 4.607 | 1.00 | 0.36 |
| MOTA | 1188 | C | GLY | 80 | 5.584 | -0.070 | 3.042 | 1.00 | 0.25 |
| MOTA | 1189 | 0 | GLY | 80 | 5.850 | -0.601 | 1.981 | 1.00 | 0.25 |
| ATOM | 1190 | N | LEU | 81 | 4.440 | 0.531 | 3.232 | 1.00 | 0.23 |
| ATOM | 1191 | HN | LEU | 81 | 4.246 | 0.951 | 4.096 | 1.00 | 0.25 |
| MOTA | 1192 | CA | LEU | 81 | 3.428 | 0.577 | 2.138 | 1.00 | 0.21 |
| ATOM | 1193 | HA | LEU | 81 | 3.259 | -0.417 | 1.761 | 1.00 | 0.22 |
| MOTA | 1194 | CB | LEU | 81 | 2.123 | 1.164 | 2.692 | 1.00 | 0.24 |

| MOTA | 1195 | HB1 L | EU | 81 | 1.587 | 1.658 | 1.896 | 1.00 | 0.25 |
|------|------|--------|-----|----------|---------|-----------|--------|------|-------|
| MCTA | 1196 | HB2 L | EU | 81 | 2.356 | 1.881 | 3.465 | 1.00 | 0.29 |
| MOTA | 1197 | CG L | EU | 81 | 1.240 | 0.058 | 3.283 | 1.00 | 0.28 |
| MOTA | 1198 | | EU | 81 | 1.856 | -0.678 | 3.779 | 1.00 | 0.31 |
| ATOM | 1199 | | EU | 81 | | _ | | | |
| | | | | | 0.265 | 0.680 | 4.285 | 1.00 | 0.33 |
| MOTA | | HD11 L | | 81 | 0.071 | 1.706 | 4.009 | 1.00 | 1.05 |
| ATOM | | HD12 L | | 81 | 0.696 | 0.649 | 5.274 | 1.00 | 1.10 |
| MOTA | 1202 | HD13 L | LEU | 81 | -0.662 | 0.125 | 4.278 | 1.00 | 1.06 |
| MOTA | 1203 | CD2 L | LEU | 81 | 0.426 | -0.606 | 2.168 | 1.00 | 0.31 |
| MOTA | 1204 | HD21 L | LEU | 81 | 1.087 | -0.997 | 1.412 | 1.00 | 1.02 |
| MOTA | | HD22 L | | 81 | -0.233 | 0.126 | 1.724 | 1.00 | 1.09 |
| ATOM | 1206 | | LEU | 81 | -0.161 | | | | |
| | | • | | | | -1.411 | 2.584 | 1.00 | 1.06 |
| MOTA | 1207 | | LEU | 81 | 3.953 | 1.475 | 1.017 | 1.00 | 0.20 |
| MOTA | 1208 | | LEU | 81 | 3.988 | 2.679 | 1.141 | 1.00 | 0.22 |
| MOTA | 1209 | N I | LEU | 82 | 4.366 | 0.899 | -0.078 | 1.00 | 0.18 |
| MOTA | 1210 | HN I | LEU | 82 | 4.334 | -0.077 | -0.162 | 1.00 | 0.18 |
| MOTA | 1211 | CA I | LEU | 82 | 4.901 | 1.728 | -1.195 | 1.00 | 0.18 |
| MOTA | 1212 | | LEU | 82 | 5.519 | 2.520 | -0.799 | 1.00 | 0.19 |
| ATOM | 1213 | | LEU | 82 | 5.728 | 0.840 | -2.128 | | |
| ATOM | 1214 | | LEU | 82 | | | | 1.00 | 0.18 |
| | | | | | 6.235 | 1.457 | -2.854 | 1.00 | 0.20 |
| MOTA | 1215 | | PEU | 82 | 5.071 | 0.151 | -2.640 | 1.00 | 0.20 |
| MOTA | 1216 | | LEU | 82 | 6.763 | 0.050 | -1.323 | 1.00 | 0.18 |
| MOTA | 1217 | HG I | LEU | 82 | 6.262 | -0.523 | -0.556 | 1.00 | 0.22 |
| MOTA | 1218 | CD1 I | LEU | 82 | 7.513 | -0.898 | -2.259 | 1.00 | 0.17 |
| MOTA | 1219 | HD11 I | LEU | 82 | 8.102 | -0.321 | -2.957 | 1.00 | 0.97 |
| MOTA | 1220 | HD12 I | LEU | 82 | 6.802 | -1.503 | -2.802 | 1.00 | 0.95 |
| ATOM | | HD13 | | 82 | 8.163 | -1.537 | -1.681 | | |
| MOTA | 1222 | CD2 I | | 00 | | | | 1.00 | 0.98 |
| | | | | | 7.764 | 1.010 | -0.675 | 1.00 | 0.23 |
| MOTA | | HD21 I | | 82 | 8.019 | 1.790 | -1.375 | 1.00 | 1.03 |
| ATOM | | | | 82 | 8.657 | 0.466 | -0.403 | 1.00 | 1.07 |
| MOTA | | HD23 I | LEU | 82 | 7.326 | 1.447 | 0.209 | 1.00 | 1.02 |
| MOTA | 1226 | CI | LEU | 82 | 3.740 | 2.329 | -1.986 | 1.00 | 0.19 |
| MOTA | 1227 | 0 1 | LEU | 82 | . 3.882 | 3.341 | -2.646 | 1.00 | 0.21 |
| ATOM | 1228 | | ALA | 83 | 2.594 | 1.711 | -1.919 | 1.00 | 0.21 |
| MOTA | 1229 | | ALA | 83 | | _ | | | |
| | | | | | 2.512 | 0.899 | -1.376 | 1.00 | 0.24 |
| MOTA | 1230 | | ALA | 83 | 1.410 | 2.225 | -2.662 | 1.00 | 0.22 |
| MOTA | 1231 | | ALA | 83 | 1.217 | 3.251 | -2.381 | 1.00 | 0.22 |
| MOTA | 1232 | CB 2 | ALA | 83 | 1.668 | 2.140 | -4.171 | 1.00 | 0.23 |
| MOTA | 1233 | HB1 | ALA | 83 | 2.522 | 2.746 | -4.429 | 1.00 | 0.98 |
| MOTA | 1234 | HB2 | ALA | 83 | 0.801 | 2.497 | -4.705 | 1.00 | 1.00 |
| MOTA | 1235 | _ | ALA | 83 | 1.860 | 1.113 | -4.445 | 1.00 | 1.05 |
| ATOM | 1236 | | ALA | 83 | 0.204 | 1.350 | -2.317 | | |
| MOTA | 1237 | | ALA | | | _ | | 1.00 | 0.27 |
| | | | | 83 | 0.342 | 0.301 | -1.720 | 1.00 | 0.36 |
| ATOM | 1238 | | HIS | 84 | -0.976 | 1.762 | -2.686 | 1.00 | 0.24 |
| MOTA | 1239 | | HIS | 84 | -1.075 | 2.609 | -3.170 | 1.00 | 0.20 |
| MOTA | 1240 | CA 1 | HIS | 84 | -2.173 | 0.933 | -2.370 | 1.00 | 0.30 |
| ATOM | 1241 | HA I | HIS | 84 | -1.940 | -0.108 | -2.542 | 1.00 | 0.36 |
| MOTA | 1242 | CB 1 | HIS | 84 | -2.562 | 1.127 | -0.903 | 1.00 | 0.40 |
| ATOM | 1243 | _ | HIS | 84 | -1.695 | 0.965 | -0.278 | 1.00 | 0.48 |
| ATOM | 1244 | _ | HIS | 84 | -3.332 | 0.419 | -0.638 | 1.00 | |
| ATOM | 1245 | | HIS | 84 | | _ | | | 0.45 |
| ATOM | 1246 | | | | -3.074 | 2.525 | -0.692 | 1.00 | 0.44 |
| _ | | | HIS | 84 | -4.384 | 2.781 | -0.321 | 1.00 | 1.32 |
| ATOM | 1247 | HD1 | | 84 | -5.084 | 2.112 | -0.169 | 1.00 | 2.02 |
| ATOM | 1248 | CD2 | | 84 | -2.465 | 3.752 | -0.788 | 1.00 | 0.74 |
| MOTA | 1249 | HD2 | | 84 | -1.432 | 3.915 | -1.060 | 1.00 | 1.58 |
| MOTA | 1250 | CE1 | HIS | 84 | -4.521 | 4.114 | -0.208 | 1.00 | 1.21 |
| MOTA | 1251 | HE1 | HIS | 84 | -5.441 | 4.606 | 0.071 | 1.00 | 1.87 |
| MOTA | 1252 | NE2 | HIS | 84 | -3.381 | 4.754 | -0.482 | 1.00 | 0.53 |
| MOTA | 1253 | C | HIS | 84 | -3.337 | 1.343 | -3.274 | 1.00 | 0.25 |
| ATOM | 1254 | | HIS | 84 | -3.347 | 2.417 | -3.843 | 1.00 | |
| ATOM | 1255 | | ALA | 85 | -4.313 | | | | 0.23 |
| ATOM | 1256 | | | | | 0.489 | -3.417 | 1.00 | 0.27 |
| | | | ALA | 85 | -4.279 | -0.374 | -2.954 | 1.00 | 0.34 |
| MOTA | 1257 | | ALA | 85 | -5.474 | 0.817 | -4.291 | 1.00 | 0.24 |
| MOTA | 1258 | | ALA | 85 | -5.582 | 1.890 | -4.364 | 1.00 | 0.22 |
| MOTA | 1259 | CB . | ALA | 85 | -5.236 | 0.231 | -5.685 | 1.00 | 0.25 |
| MOTA | 1260 | HB1 | ALA | 85 | -5.079 | -0.835 | -5.605 | 1.00 | 1.05 |
| ATOM | 1261 | HB2 | | 85 | -4.364 | 0.690 | -6.126 | 1.00 | 1.05 |
| ATOM | 1262 | нвз | | 85 | -6.097 | 0.420 | -6.308 | | |
| ATOM | 1263 | | ALA | 85 | | | | 1.00 | 1.06 |
| MOTA | 1264 | | | | -6.748 | 0.210 | -3.698 | 1.00 | 0.26 |
| | | | ALA | 85 86 | -6.694 | -0.611 | -2.804 | 1.00 | 0.33 |
| ATOM | 1265 | | PHE | 86 | -7.892 | 0.605 | -4.198 | 1.00 | 0.28 |
| ATOM | 1266 | | PHE | 86 | -7.905 | 1.264 | -4.922 | 1.00 | 0.31 |
| MOTA | 1267 | | PHE | 86 | -9.179 | 0.053 | -3.677 | 1.00 | 0.34 |
| MOTA | 1268 | HA | PHE | 86 | -9.000 | -0.443 | -2.737 | 1.00 | 0.39 |
| MOTA | 1269 | | PHE | 86 | -10.170 | 1.205 | -3.471 | 1.00 | 0.36 |
| MOTA | 1270 | | PHE | 86 | -11.177 | 0.821 | -3.459 | 1.00 | 0.42 |
| MOTA | 1271 | HB2 | | 86 | -10.068 | 1.913 | -4.279 | 1.00 | 0.33 |
| | | | | | | ~ · 3 T 3 | -4.413 | 1.00 | U. J. |

| | 1 222 | | | 0.0 | | | | | |
|-------------|-------|-------|-----|------|-----------------|--------|------------------|------|------|
| MOTA | 1272 | | PHE | 86 | -9.877 | 1.896 | -2.159 | 1.00 | 0.39 |
| MOTA | 1273 | CD1 | PHE | 86 | -8.784 | 2.764 | -2.050 | 1.00 | 0.46 |
| MOTA | 1274 | HD1 | PHE | 86 | -8.146 | 2.939 | -2.903 | 1.00 | 0.67 |
| MOTA | 1275 | CD2 | PHE | 86 | -10.703 | 1.670 | -1.051 | 1.00 | 0.67 |
| ATOM | 1276 | | PHE | 86 | | | | | |
| | | | | | -11.546 | 1.001 | -1.133 | 1.00 | 0.91 |
| ATOM | 1277 | | PHE | 86 | -8.516 | 3.406 | -0.835 | 1.00 | 0.50 |
| MOTA | 1278 | HE1 | PHE | 86 | - 7. 673 | 4.075 | -0.751 | 1.00 | 0.69 |
| ATOM | 1279 | CE2 | PHE | 86 | -10.435 | 2.311 | 0.165 | 1.00 | 0.74 |
| MOTA | 1280 | HE2 | PHE | 86 | -11.071 | 2.136 | 1.020 | 1.00 | 1.02 |
| | | | | | | | | | _ |
| MOTA | 1281 | CZ | PHE | 86 | -9.342 | 3.179 | 0.273 | 1.00 | 0.54 |
| MOTA | 1282 | HZ | PHE | 86 | -9.135 | 3.674 | 1.211 | 1.00 | 0.62 |
| MOTA | 1283 | С | PHE | 86 | -9.746 | -0.940 | -4.710 | 1.00 | 0.36 |
| MOTA | 1284 | 0 | PHE | 86 | -9.480 | -0.812 | -5.889 | 1.00 | 0.34 |
| MOTA | 1285 | N | PRO | 87 | | | | | |
| | | | | | -10.516 | -1.926 | -4.293 | 1.00 | 0.43 |
| MOTA | 1286 | CA | PRO | 87 | -11.082 | -2.914 | -5.257 | 1.00 | 0.46 |
| MOTA | 1287 | HA | PRO | 87 | -10.296 | -3.524 | -5.665 | 1.00 | 0.53 |
| ATOM . | 1288 | CB | PRO | 87 | -11.990 | -3.770 | -4.370 | 1.00 | 0.60 |
| MOTA | 1289 | HB1 | PRO | 87 | -11.644 | -4.792 | -4.377 | 1.00 | 0.69 |
| | 1290 | | | | _ | _ | | | |
| ATOM | | HB2 | PRO | 87 | -13.004 | -3.727 | -4.742 | 1.00 | 0.73 |
| MOTA | 1291 | CG | PRO | 87 | -11.943 | -3,225 | -2.937 | 1.00 | 0.58 |
| MOTA | 1292 | HG1 | PRO | 87 | -11.694 | -4.022 | -2.253 | 1.00 | 0.61 |
| MOTA | 1293 | HG2 | PRO | 87 | -12.905 | -2.808 | -2.676 | 1.00 | 0.66 |
| ATOM | 1294 | | PRO | 87 | | | | | _ |
| | | CD | _ | | -10.872 | -2.135 | -2.861 | 1.00 | 0.50 |
| MOTA | 1295 | HD2 | PRO | 87 | -11.277 | -1.235 | -2.421 | 1.00 | 0.50 |
| MOTA | 1296 | HD1 | PRO | 87 | -10.014 | -2.484 | -2.309 | 1.00 | 0.52 |
| ATOM | 1297 | C | PRO | 87 | -11.895 | -2.246 | -6.379 | 1.00 | 0.40 |
| MOTA | 1298 | 0 | PRO | 87 | -12.221 | -1.078 | -6.299 | 1.00 | 0.42 |
| ATOM | 1299 | Ň | | | | | | | _ |
| | | - | PRO | 88 | -12.221 | -2.981 | -7.419 | 1.00 | 0.44 |
| MOTA | 1300 | CA | PRO | 88 | -13.007 | -2.416 | -8.554 | 1.00 | 0.48 |
| MOTA | 1301 | HA | PRO | 88 | -12.443 | -1.645 | -9.053 | 1.00 | 0.52 |
| MOTA | 1302 | CB | PRO | 88 | -13.163 | -3.622 | -9.488 | 1.00 | 0.61 |
| ATOM | 1303 | HB1 | PRO | | | | • • - • - | | |
| | | | | 88 | -12.604 | | -10.395 | 1.00 | 0.83 |
| MOTA | 1304 | HB2 | PRO | 88 | -14.204 | -3.772 | -9.728 | 1.00 | 0.74 |
| MOTA | 1305 | CG | PRO | 88 | -12.609 | -4.863 | -8.781 | 1.00 | 0.57 |
| ATOM | 1306 | HG1 | PRO | 88 | -11.945 | -5.395 | -9.446 | 1.00 | 0.71 |
| ATOM | 1307 | HG2 | PRO | 88 | -13.425 | -5.508 | -8.488 | | _ |
| | | _ | | | | | | 1.00 | 0.64 |
| MOTA | 1308 | CD | PRO | 88 | -11.835 | -4.413 | -7.540 | 1.00 | 0.56 |
| MOTA | 1309 | HD2 | PRO | 88 | -12.146 | -4,977 | -6.671 | 1.00 | 0.62 |
| MOTA | 1310 | - HD1 | PRO | 88 | -10.773 | -4.503 | -7.702 | 1.00 | 0.65 |
| ATOM | 1311 | C | PRO | 88 | -14.372 | -1.873 | -8.109 | 1.00 | 0.47 |
| ATOM | 1312 | · _ | PRO | | | | _ | | |
| | | 0 | | 88 | -15.380 | -2.551 | -8.172 | 1.00 | 0.88 |
| MOTA | 1313 | N | GLY | 89 | -14.400 | -0.647 | -7.661 | 1.00 | 0.63 |
| ATOM | 1314 | HN | GLY | 89 | -13.571 | -0.129 | -7.626 | 1.00 | 1.01 |
| ATOM | 1315 | CA | GLY | 89 | -15.681 | -0.026 | -7.209 | 1.00 | 0.65 |
| ATOM | 1316 | HA1 | | 89 | -15.536 | 0.422 | -6.239 | 1.00 | 0.62 |
| ATOM | 1317 | HA2 | | | | | | | |
| | | | | 89 | -16.455 | -0.778 | -7.148 | 1.00 | 0.78 |
| ATOM | 1318 | C | GLY | 89 | -16.092 | 1.057 | ~8.210 | 1.00 | 0.74 |
| MOTA | 1319 | 0 | GLY | 89 | -15.541 | 1.151 | ~9.289 | 1.00 | 0.84 |
| ATOM | 1320 | N | PRO | 90 | -17.044 | 1.878 | -7.852 | 1.00 | 0.95 |
| ATOM | 1321 | CA | PRO | 90 | -17.499 | 2.973 | -8.750 | 1.00 | 1.19 |
| ATOM | 1322 | | | | | | | | |
| | | HA | PRO | 90 | -17.819 | 2.565 | -9.697 | 1.00 | 1.37 |
| MOTA | 1323 | CB | PRO | 90 | -18.720 | 3.532 | -7.990 | 1.00 | 1.55 |
| ATOM | 1324 | HB1 | PRO | 90 | -19.602 | 3.432 | -8.605 | 1.00 | 1.85 |
| MOTA | 1325 | HB2 | PRO | 90 | -18.572 | 4.567 | -7.740 | 1.00 | 1.74 |
| MOTA | 1326 | CG | PRO | 90 | -18.913 | 2.724 | -6.702 | 1.00 | 1.46 |
| MOTA | 1327 | HG1 | | 90 | -19.828 | 2.155 | | | |
| ATOM | 1328 | | | | | | -6.763 | 1.00 | 1.60 |
| | | HG2 | | 90 | -18.959 | 3.396 | -5.857 | 1.00 | 1.57 |
| MOTA | 1329 | CD | PRO | 90 | -17.729 | 1.769 | -6.539 | 1.00 | 1.17 |
| ATOM | 1330 | HD2 | PRO | 90 | -17.083 | 2.099 | -5.736 | 1.00 | 1.17 |
| MOTA | 1331 | HD1 | PRO | 90 | -18.067 | 0.759 | -6.375 | 1.00 | 1.28 |
| ATOM | 1332 | C | PRO | 90 | -16.375 | 4.011 | -8.972 | | |
| ATOM | 1333 | | | | | | | 1.00 | 1.14 |
| | | 0 | PRO | 90 | -15.269 | 3.649 | -9.320 | 1.00 | 1.53 |
| MOTA | 1334 | N | asn | 91 | -16.624 | 5,282 | -8.790 | 1.00 | 1.17 |
| MOTA | 1335 | HN | asn | 91 | -17.514 | 5.578 | -8.517 | 1.00 | 1.40 |
| MOTA | 1336 | CA | ASN | 91 | -15.541 | 6.286 | -9.008 | 1.00 | 1.38 |
| ATOM | 1337 | HA | ASN | 91 | -15.147 | | -10.005 | | |
| ATOM | 1338 | | | | | | | 1.00 | 1.58 |
| | | CB | ASN | 91 | -16.116 | 7.700 | -8.857 | 1.00 | 1.87 |
| MOTA | 1339 | HB1 | | 91 | -15.336 | 8.372 | -8.532 | 1.00 | 2.33 |
| MOTA | 1340 | HB2 | ASN | 91 | -16.908 | 7.686 | | | 1.96 |
| ATOM | 1341 | CG | ASN | 91 | -16.678 | | -10.197 | 1.00 | 2.69 |
| MOTA | 1342 | | ASN | 91 | -16.132 | | | | |
| | | | | | | 7.890 | | 1.00 | 3.20 |
| MOTA | 1343 | | ASN | · 91 | -17.748 | | -10.212 | 1.00 | 3.47 |
| MOTA | 1344 | | ASN | 91 | -18.186 | 9.176 | -9.370 | 1.00 | 3.59 |
| MOTA | 1345 | HD22 | ASN | 91 | -18.112 | 9,249 | | 1.00 | 4.20 |
| MOTA | 1346 | С | ASN | 91 | -14.404 | 6.098 | | 1.00 | 1.15 |
| MOTA | 1347 | Ö | ASN | 91 | -13.242 | 6.135 | | 1.00 | 1.26 |
| ATOM | 1348 | N | TYR | 92 | -14 719 | 5 924 | -6.344 -£ 735 | 1 00 | 1 01 |
| | | | | | ,, 4 | /4 | / \ | | |

| ATOM | 1349 | HN | TYR | 92 | -15 660 | 5 016 | 6 460 | | 1 00 |
|--------|------|-----|-----|-----------|----------------|--------|--------|------|-------|
| MOTA | 1350 | CA | TYR | | -15.660 | 5.916 | -6.462 | 1.00 | 1.08 |
| | 1351 | | | 92 | -13.639 | 5.768 | -5.711 | 1.00 | 0.97 |
| MOTA | _ | HA | TYR | 92 | -12.994 | 6.632 | -5.739 | 1.00 | 1.14 |
| MOTA | 1352 | CB | TYR | 92 | -14.262 | 5.652 | -4.319 | 1.00 | 1.09 |
| MOTA | 1353 | HB1 | TYR | 92 | -13.543 | 5.214 | -3.643 | 1.00 | 1.62 |
| ATOM . | 1354 | HB2 | TYR | 92 | -15.135 | 5.020 | -4.369 | 1.00 | 1.45 |
| MOTA | 1355 | CG | TYR | 92 | -14.656 | 7.018 | -3.810 | 1.00 | 1.52 |
| MOTA | 1356 | CD1 | TYR | 92 | -13.672 | 7.979 | -3.549 | 1.00 | 2.14 |
| ATOM | 1357 | HD1 | TYR | 92 | -12.631 | 7.747 | -3.719 | 1.00 | 2.46 |
| MOTA | 1358 | CD2 | TYR | 92 | -16.006 | 7.320 | -3.588 | 1.00 | 2.44 |
| ATOM | 1359 | HD2 | TYR | 92 | -16.766 | 6.580 | -3.789 | 1.00 | 2.86 |
| MOTA | 1360 | CE1 | TYR | 92 | -14.037 | 9.241 | -3.066 | 1.00 | 3.06 |
| MOTA | 1361 | HE1 | TYR | 92 | -13.278 | 9.982 | -2.865 | 1.00 | 3.78 |
| ATOM | 1362 | CE2 | TYR | 92 | -16.370 | 8.582 | -3.107 | | |
| ATOM | 1363 | HE2 | TYR | 92 | -17.411 | _ | | 1.00 | 3.33 |
| ATOM | 1364 | CZ | TYR | 92 | -15.386 | 8.815 | -2.936 | 1.00 | 4.19 |
| MOTA | 1365 | OH | TYR | 92 | · - | 9.542 | -2.846 | 1.00 | 3.50 |
| MOTA | 1365 | | | | -15.746 | 10.786 | -2.368 | 1.00 | 4.57 |
| | | НН | TYR | 92 | -15.602 | 10.791 | -1.419 | 1.00 | 4.91 |
| MOTA | 1367 | C | TYR | 92 | -12.808 | 4.508 | -5.966 | 1.00 | 0.78 |
| ATOM | 1368 | 0 | TYR | 92 | -11.605 | 4.506 | -5.798 | 1.00 | 0.81 |
| MOTA | 1369 | N | GLY | 93 | -13.436 | 3.430 | -6.337 | 1.00 | 0.64 |
| MOTA | 1370 | HN | GLY | 93 | -14.410 | 3.441 | -6.445 | 1.00 | 0.70 |
| ATOM | 1371 | CA | GLY | 93 | -12.674 | 2.170 | -6.560 | 1.00 | 0.51 |
| ATOM | 1372 | HA1 | GLY | 93 | -13.366 | 1.366 | -6.740 | 1.00 | 0.51 |
| MOTA | 1373 | HA2 | GLY | 93 | -12.090 | 1.947 | -5.678 | 1.00 | 0.51 |
| MOTA | 1374 | Ç | GLY | 93 | -11.739 | 2.310 | -7.761 | 1.00 | 0.49 |
| ATOM | 1375 | 0 | GLY | 93 | -11.832 | 3.242 | -8.534 | 1.00 | 0.61 |
| MOTA | 1376 | N | GLY | 94 | -10.844 | 1.373 | -7.923 | 1.00 | |
| MOTA | 1377 | HN | GLY | 94 | -10.799 | 0.627 | | | 0.45 |
| ATOM | 1378 | CA | GLY | 94 | | | -7.288 | 1.00 | 0.44 |
| ATOM | 1379 | HA1 | | | -9.902 | 1.420 | -9.075 | 1.00 | 0.55. |
| MOTA | 1380 | HA2 | | 94 | -10.459 | 1.569 | -9.988 | 1.00 | 0.63 |
| ATOM | | | | 94 | -9.363 | 0.485 | -9.133 | 1.00 | 0.58 |
| | 1381 | C | GLY | 94 | -8.905 | 2.569 | -8.901 | 1.00 | 0.60 |
| MOTA. | 1382 | 0 | GLY | 94 | -8.109 | 2.838 | -9.772 | 1.00 | 1.14 |
| MOTA | 1383 | N | ASP | 95 | -8.933 | 3.252 | -7.790 | 1.00 | 0.24 |
| MOTA | 1384 | HN | ASP | 95 | -9.581 | 3.028 | -7.089 | 1.00 | 0.52 |
| MOTA | 1385 | CA | ASP | 95 | -7.976 | 4.382 | -7.597 | 1.00 | 0.24 |
| ATOM | 1386 | HA | ASP | 95 | -7.888 | 4.939 | -8.518 | 1.00 | 0.28 |
| MOTA | 1387 | CB | ASP | 95 | -8.493 | 5.303 | -6.491 | 1.00 | 0.26 |
| MOTA | 1388 | HB1 | ASP | 95 | -9.500 | 5.617 | -6.724 | 1.00 | 0.28 |
| ATOM | 1389 | HB2 | _ | 95 | -7.853 | 6.170 | -6.415 | 1.00 | |
| MOTA | 1390 | CG | ASP | 95 | -8.494 | 4.549 | | _ | 0.30 |
| MOTA | 1391 | - | ASP | 95 | -8.543 | | -5.162 | 1.00 | 0.28 |
| ATOM | 1392 | | ASP | 95 | | 5.200 | -4.132 | 1.00 | 1.08 |
| ATOM | 1393 | C | | | -8.440 | 3.331 | -5.198 | 1.00 | 1.14 |
| ATOM | | | ASP | 95 05 | -6.605 | 3.827 | -7.202 | 1.00 | 0.23 |
| | 1394 | 0 | ASP | 95 | -6.479 | 2.683 | -6.815 | 1.00 | 0.24 |
| MOTA | 1395 | N | ALA | 96 | -5.573 | 4.626 | -7.297 | 1.00 | 0.23 |
| MOTA | 1396 | HN | ALA | 96 | -5.692 | 5.546 | -7.614 | 1.00 | 0.23 |
| MOTA | 1397 | CA | ALA | 96 | -4.215 | 4.131 | -6.926 | 1.00 | 0.25 |
| MOTA | 1398 | HA | ALA | 96 | -4.307 | 3:360 | -6.175 | 1.00 | 0.25 |
| MOTA | 1399 | CB | ALA | 96 | -3.527 | 3.553 | -8.164 | 1.00 | 0.30 |
| MOTA | 1400 | HB1 | ALA | 96 | -2.528 | 3.236 | -7.905 | 1.00 | 1.08 |
| MOTA | 1401 | HB2 | ALA | 96 | -3.476 | 4.309 | -8.934 | 1.00 | 1.08 |
| MOTA | 1402 | HB3 | ALA | 96 | -4.090 | 2.706 | -8.528 | 1.00 | 1.03 |
| MOTA | 1403 | С | ALA | 96 | -3.375 | 5.284 | -6.372 | 1.00 | 0.25 |
| MOTA | 1404 | 0 | ALA | 96 | -3.222 | 6.313 | -7.005 | 1.00 | |
| ATOM | 1405 | N | HXS | 97 | -2.831 | 5.113 | -5.192 | | 0.29 |
| ATOM | 1406 | HN | HXS | 97 | -2.976 | | _ | 1.00 | 0.25 |
| MOTA | 1407 | CA | HXS | 97 | -1.996 | 4,271 | -4.710 | 1.00 | 0.28 |
| ATOM | 1408 | HA | HXS | 97 | | 6.187 | -4.574 | 1.00 | 0.27 |
| ATOM | 1409 | CB | | | -2.010 | 7.068 | -5.198 | 1.00 | 0.28 |
| ATOM | 1410 | | HXS | 97 22 | -2.564 | 6.537 | -3.197 | 1.00 | 0.33 |
| | | HB1 | | 97 | -1.969 | 7.319 | ~2.750 | 1.00 | 0.44 |
| ATOM | 1411 | HB2 | | 97 | -2.540 | 5.661 | -2.566 | 1.00 | 0.39 |
| ATOM | 1412 | CG | HXS | 97 | -3.983 | 7.009 | -3.349 | 1.00 | 0.37 |
| MOTA | 1413 | | HXS | 97 | -4.697 | 7.052 | -2.163 | 1.00 | 0.80 |
| MOTA | 1414 | | HXS | 97 | -4.783 | 7.420 | -4.384 | 1.00 | 0.55 |
| MOTA | 1415 | | HXS | 97 | -4.517 | 7.497 | -5.428 | 1.00 | 0.94 |
| MOTA | 1416 | | HXS | 97 | -5.918 | 7.487 | -2.498 | 1.00 | 0.86 |
| MOTA | 1417 | | HXS | 97 | -6.724 | 7.632 | -1.795 | 1.00 | 1.24 |
| MOTA | 1418 | | HXS | 97 | -6.018 | 7.722 | -3.819 | | |
| MOTA | 1419 | HE2 | | 97 | -6.812 | 8.044 | | 1.00 | 0.59 |
| MOTA | 1420 | C | HXS | 97 | -0.552 | 5.700 | -4.294 | 1.00 | 0.72 |
| ATOM | 1421 | Õ | HXS | 97 | | | -4.420 | 1.00 | 0.26 |
| ATOM | 1422 | N | PHE | | -0.299 | 4.525 | -4.237 | 1.00 | 0.39 |
| ATOM | 1423 | | | 98 | 0.391 | 6.604 | -4.496 | 1.00 | 0.18 |
| ATOM | 1424 | HN | PHE | 98 | 0.147 | 7.540 | -4.648 | 1.00 | 0.23 |
| MOTA | | CA | PHE | 98 | 1.832 | 6.230 | -4.360 | 1.00 | 0.17 |
| UM | 1425 | HA | PHE | 98 | 1.921 | 5.190 | -4.085 | 1.00 | 0.18 |

| ATOM | 1426 | ÇВ | PHE | 98 | 2.543 | 6.472 | -5.691 | 1.00 | 0.18 |
|--------------|--------------|-----------|------------|--------------|------------------|------------------|------------------|------|--------------|
| MOTA | 1427 | HB1 | PHE | 98 | 3.611 | 6.464 | -5.536 | 1.00 | 0.21 |
| ATOM | 1428 | | PHE | 98 | 2.243 | 7.431 | -6.085 | 1.00 | 0.20 |
| ATOM | 1429 | | PHE | 98 | 2.169 | 5.391 | -6.674 | 1.00 | 0.19 |
| MOTA MOTA | 1430 1431 | | PHE PHE | 98 98 | 3.114 4.110 | 4.428 4.456 | -7.048 | 1.00 | 0.22 |
| ATOM | 1432 | _ | PHE | 98 | 0.880 | 5.355 | -6.631 -7.214 | 1.00 | 0.25 0.22 |
| ATOM | 1433 | | PHE | 98 | 0.151 | 6.098 | -6.924 | 1.00 | 0.24 |
| ATOM | 1434 | | PHE | 98 | 2.768 | 3.429 | -7.963 | 1.00 | 0.25 |
| MOTA | 1435 | | PHE | 98 | 3.496 | 2.685 | -8.252 | 1.00 | 0.29 |
| MOTA MOTA | 1436 1437 | | PHE | 98 | 0.533 | 4.355 | -8.127 | 1.00 | 0.26 |
| ATOM | 1438 | HE2 CZ | PHE PHE | 98 98 | -0.462 1.478 | 4.327 3.392 | -8.542 -8.503 | 1.00 | 0.31 |
| ATOM | 1439 | HZ | PHE | 98 | 1.214 | 2.622 | -9.211 | 1.00 | 0.26 0.30 |
| MOTA | 1440 | С | PHE | 98 | 2.487 | 7.104 | -3.286 | 1.00 | 0.17 |
| MOTA | 1441 | 0 | PHE | 98 | 2.081 | 8.226 | -3.058 | 1.00 | 0.19 |
| MOTA MOTA | 1442 1443 | N | ASP | 99 | 3.498 | 6.604 | -2.625 | 1.00 | 0.19 |
| ATOM | 1444 | HN CA | ASP ASP | 99 99 | 3.813 4.167 | 5.693 7.424 | -2.820 -1.570 | 1.00 | 0.22 |
| ATOM | 1445 | HA | ASP | 99 | 3.421 | 7.956 | -0.998 | 1.00 | 0.20 0.20 |
| MOTA | 1446 | CB | ASP | 99 | 4.973 | 6.516 | -0.638 | 1.00 | 0.25 |
| MOTA | 1447 | | | 99 | 5.567 | 7.122 | 0.029 | 1.00 | 0.28 |
| MOTA MOTA | 1448 1449 | HB2 CG | ASP | 99 | 5.624 | 5.884 | -1.226 | 1.00 | 0.30 |
| MOTA | 1450 | | ASP ASP | 99 99 | 4.023 2.838 | 5.646 5.680 | 0.180 -0.100 | 1.00 | 0.41 |
| MOTA | 1451 | _ | ASP | 99 | 4.497 | 4.968 | 1.079 | 1.00 | 0.89 0.27 |
| MOTA | 1452 | C | ASP | 99 | 5.123 | 8.426 | -2.224 | 1.00 | 0.21 |
| MOTA | 1453 | 0 | ASP | 99 | 6.020 | 8.054 | -2.954 | 1.00 | 0.25 |
| ATOM ATOM | 1454 1455 | N HN | ASP | 100 | 4.946 | 9.694 | -1.962 | 1.00 | 0.23 |
| ATOM | 1456 | CA | ASP ASP | 100 100 | 4.222 5.857 | 9.976 10.710 | -1.365 | 1.00 | 0.23 |
| ATOM | 1457 | HA | ASP | 100 | 6.169 | 10.710 | -2.565 -3.545 | 1.00 | 0.29 0.31 |
| MOTA | 1458 | CB | ASP | 100 | 5.127 | 12.049 | -2.684 | 1.00 | 0.34 |
| MOTA | 1459 | HB1 | | 100 | 5.130 | 12.544 | -1.727 | 1.00 | 0.34 |
| MOTA MOTA | 1460 | | ASP | 100 | 4.109 | 11.879 | -2.999 | 1.00 | 0.34 |
| ATOM | 1461 1462 | CG | ASP ASP | 100 100 | 5.844 | 12.929 | -3.710 | 1.00 | 0.43 |
| ATOM | 1463 | | ASP | 100 | 5.240 6.984 | 13.887 12.630 | -4.164 -4.025 | 1.00 | 1.21 1.12 |
| MOTA | 1464 | C | ASP | 100 | 7.085 | 10.885 | -1.667 | 1.00 | 0.30 |
| ATOM | 1465 | 0 | ASP | 100 | 8.032 | 11.559 | -2.018 | 1.00 | 0.32 |
| MOTA MOTA | 1466 | N | ASP | 101 | 7.074 | 10.280 | -0.510 | 1.00 | 0.31 |
| ATOM | 1467 1468 | HN CA | ASP ASP | 101 101 | 6.298 | 9.741 | -0.249 | 1.00 | 0.32 |
| ATOM | 1469 | HA | ASP | 101 | 8.236 8.647 | 10.407 11.403 | 0.415 0.345 | 1.00 | 0.33 0.36 |
| MOTA | 1470 | CB | ASP | 101 | 7.778 | 10.142 | 1.851 | 1.00 | 0.39 |
| MOTA | 1471 | HB1 | | 101 | 8.641 | 10.060 | 2.495 | 1.00 | 0.41 |
| MOTA MOTA | 1472 1473 | | ASP | 101 | 7.216 | 9.220 | 1.884 | 1.00 | 0.39. |
| ATOM | 1474 | CG OD1 | ASP ASP | 101 101 | 6.896 7.027 | 11.296 12.380 | 2.330 | 1.00 | 0.45 |
| ATOM | 1475 | | | 101 | 6.104 | 11.076 | 1.786 3.231 | 1.00 | 1.25 1.09 |
| MOTA | 1476 | С | ASP | 101 | 9.304 | 9.385 | 0.028 | 1.00 | 0.30 |
| ATOM | 1477 | 0 | ASP | 101 | 10.411 | 9.405 | 0.529 | 1.00 | 0.29 |
| MOTA MOTA | 1478 1479 | N HN | GLU GLU | 102 | 8.971 | 8.484 | -0.849 | 1.00 | 0.30 |
| ATOM | 1480 | CA | GLU | 102 102 | 8.068 9.950 | 8.484 7.444 | -1.230 -1.266 | 1.00 | 0.31 |
| MOTA | 1481 | HA | GLU | 102 | 10.649 | 7.263 | -0.463 | 1.00 | 0.29 0.30 |
| MOTA | 1482 | CB | GLU | 102 | 9.195 | 6.155 | -1.585 | 1.00 | 0.35 |
| ATOM ATOM | 1483 | | GLU | 102 | 9.873 | 5.437 | -2.020 | 1.00 | 0.36 |
| ATOM | 1484 1485 | HB2 CG | GLU GLU | 102 · 102 | 8.397 8.611 | 6.368 | -2.282 | 1.00 | 0.40 |
| MOTA | 1486 | HG1 | | 102 | 8.020 | 5.584 6.342 | -0.293 0.200 | 1.00 | 0.46 1.18 |
| MOTA | 1487 | HG2 | | 102 | 9.415 | 5.276 | 0.356 | 1.00 | 1.03 |
| ATOM | 1488 | CD | GLU | 102 | 7.724 | 4.381 | -0.616 | 1.00 | 0.83 |
| MOTA MOTA | 1489 | OE1 | | 102 | 7.601 | 4.060 | -1.786 | 1.00 | 1.63 |
| ATOM | 1490 1491 | OE2 C | GLU | 102 102 | 7.184 | 3.801 | 0.314 | 1.00 | 0.87 |
| MOTA | 1492 | ŏ | GLU | 102 | 10.707 10.359 | 7.917 8.910 | -2.508 -3.115 | 1.00 | 0.25 0.25 |
| ATOM | 1493 | N | THR | 103 | 11.741 | 7.213 | -2.886 | 1.00 | 0.25 |
| MOTA | 1494 | HN | THR | 103 | 12.003 | 6.416 | -2.379 | 1.00 | 0.28 |
| MOTA MOTA | 1495 1496 | CA | THR | 103 | 12.525 | 7.620 | -4.088 | 1.00 | 0.23 |
| ATOM | 1495 | HA CB | THR | 103 103 | 12.356 | 8.665 | -4.301 | 1.00 | 0.23 |
| ATOM | 1498 | HB | THR | 103 | 14.016 14.169 | 7.383 6.359 | -3.824 -3.521 | 1.00 | 0.27 0.30 |
| MOTA | 1499 | 0G1 | | 103 | 14.455 | 8.252 | -3.521 -2.789 | 1.00 | 0.30 |
| ATOM | 1500 | HG1 | THR | 103 | 15.334 | 8.564 | -3.016 | 1.00 | 0.86 |
| MOTA MOTA | 1501 1502 | HG21 | THR | 103 | 14.820 | 7.656 | -5.098 | 1.00 | 0.29 |
| | | | TUK | 103 . | 15.864 | 7 777 | -4 845 | 1 00 | 1 00 |

| MOTA | | HG22 | THR | 103 | 14.457 | 8.557 | -5.569 | 1.00 | 1.08 |
|------|------|------|------|-------|--------|--------|---------|------|------|
| MOTA | 1504 | HG23 | THR | 103 | 14.710 | 6.824 | -5.779 | 1.00 | 1.01 |
| MOTA | 1505 | C | THR | 103 | 12.083 | | -5.281 | _ | |
| ATOM | 1506 | Ö | THR | 103 | | 6.777 | | 1.00 | 0.22 |
| | | | | | 12.417 | 5.614 | -5.394 | 1.00 | 0.23 |
| MOTA | 1507 | N | TRP | 104 | 11.332 | 7.358 | -6.175 | 1.00 | 0.21 |
| MOTA | 1508 | HN | TRP | 104 | 11.076 | 8.297 | -6.063 | 1.00 | 0.23 |
| MOTA | 1509 | CA | TRP | 104 | 10.867 | 6.598 | -7.364 | | |
| MOTA | 1510 | HA | TRP | | | | | 1.00 | 0.21 |
| | | | | 104 | 10.750 | 5.556 | -7.104 | 1.00 | 0.20 |
| MOTA | 1511 | CB | TRP | 104 | 9.525 | 7.165 | -7.831 | 1.00 | 0.23 |
| ATOM | 1512 | HB1 | TRP | 104 | 9.188 | 6.623 | -8.702 | 1.00 | 0.24 |
| MOTA | 1513 | HB2 | TRP | 104 | 9.641 | | | _ | |
| ATOM | 1514 | CG | | | | 8.210 | -8.078 | 1.00 | 0.25 |
| | | | TRP. | 104 | 8.520 | 7.018 | -6.731 | 1.00 | 0.24 |
| MOTA | 1515 | CD1 | | 104 | 8.098 | 8.019 | -5.924 | 1.00 | 0.31 |
| ATOM | 1516 | HD1 | TRP | 104 | 8.427 | 9.045 | -5.972 | 1.00 | |
| MOTA | 1517 | CD2 | TRP | 104 | | | | | 0.36 |
| MOTA | 1518 | | | | 7.811 | 5.821 | -6.300 | 1.00 | 0.21 |
| | | | TRP | 104 | 7.176 | 7.512 | -5.026 | 1.00 | 0.31 |
| MOTA | 1519 | | TRP | 104 | 6.718 | 8.030 | -4.331 | 1.00 | 0.36 |
| MOTA | 1520 | CE2 | TRP | 104 | 6.963 | 6.162 | -5.220 | 1.00 | 0.24 |
| MOTA | 1521 | CE3 | TRP | 104 | 7.819 | | | - | |
| MOTA | 1522 | | TRP | | | 4.486 | -6.739 | 1.00 | 0.18 |
| | | | | 104 | 8.458 | 4.198 | -7.559 | 1.00 | 0.19 |
| MOTA | 1523 | | TRP | 104 | 6.153 | 5.213 | -4.596 | 1.00 | 0.23 |
| MOTA | 1524 | HZ2 | TRP | 104 | 5.515 | 5.499 | -3.774 | 1.00 | 0.27 |
| MOTA | 1525 | CZ3 | TRP | 104 | 7.005 | | | | |
| ATOM | 1526 | HZ3 | | | | 3.527 | -6.114 | 1.00 | 0.20 |
| | | | | 104 | 7.019 | 2.504 | -6.460 | 1.00 | 0.23 |
| MOTA | 1527 | CH2 | _ | 104 | 6.173 | 3.891 | -5.045 | 1.00 | 0.21 |
| MOTA | 1528 | HH2 | TRP | 104 | 5.548 | 3.150 | -4.568 | 1.00 | 0.23 |
| MOTA | 1529 | С | TRP | 104 | 11.911 | 6.732 | | | |
| ATOM | 1530 | Ŏ | TRP | | | | -8.474 | 1.00 | 0.21 |
| | | | | 104 | 12.276 | 7.824 | -8.864 | 1.00 | 0.24 |
| MOTA | 1531 | N | THR | 105 | 12.403 | 5.630 | -8.973 | 1.00 | 0.20 |
| ATOM | 1532 | HN | THR | 105 | 12.098 | 4.763 | | 1.00 | 0.19 |
| MOTA | 1533 | CA | THR | 105 | 13.437 | | | | |
| MOTA | 1534 | HA | THR | _ | | | -10.048 | 1.00 | 0.21 |
| | | | | 105 | 13.415 | 6.652 | -10.525 | 1.00 | 0.24 |
| ATOM | 1535 | CB | THR | 105 | 14.817 | 5.459 | -9.428 | 1.00 | 0.21 |
| MOTA | 1536 | HB | THR | 105 | 15.018 | 6.233 | | 1.00 | |
| ATOM | 1537 | OG1 | THR | 105 | 15.806 | | | | 0.21 |
| MOTA | 1538 | | THR | | | | -10.447 | 1.00 | 0.24 |
| | | | | 105 | 15.882 | 6.404 | -10.752 | 1.00 | 0.86 |
| MOTA | 1539 | | THR | . 105 | 14.846 | 4.101 | -8.729 | 1.00 | 0.21 |
| MOTA | | HG21 | THR | 105 | 15.178 | 4.233 | | 1.00 | |
| ATOM | 1541 | HG22 | THR | 105 | 15.524 | | | | 1.04 |
| MOTA | | HG23 | | | | 3.442 | | 1.00 | 1.07 |
| | | | | 105 | 13.854 | 3.674 | -8.731 | 1.00 | 0.99 |
| MOTA | 1543 | C | THR | 105 | 13.166 | 4.597 | -11.087 | 1.00 | 0.23 |
| MOTA | 1544 | 0 | THR | 105 | 12.521 | | -10.808 | | |
| ATOM | 1545 | N | SER | 106 | | | | 1.00 | 0.23 |
| MOTA | 1546 | HN | | | 13.668 | | -12.282 | 1.00 | 0.26 |
| | | | SER | 106 | 14.194 | 5.572 | -12.480 | 1.00 | 0.29 |
| MOTA | 1547 | CA | SER | 106 | 13.454 | 3.739 | -13.337 | 1.00 | 0.29 |
| MOTA | 1548 | HA | SER | 106 | 12.570 | | -13.111 | 1.00 | 0.30 |
| ATOM | 1549 | CB | SER | 106 | 13.290 | | -14.695 | | |
| MOTA | 1550 | HB1 | | 106 | | | | 1.00 | 0.35 |
| MOTA | | | | | 14.249 | | -15.193 | 1.00 | 1.09 |
| | 1551 | HB2 | | 106 | 12.916 | 5.424 | -14.554 | 1.00 | 0.96 |
| MOTA | 1552 | OG | SER | 106 | 12.365 | | -15.483 | 1.00 | 1.44 |
| MOTA | 1553 | HG | SER | 106 | 11.671 | | -15.766 | | |
| ATOM | 1554 | С | SER | 106 | | | | 1.00 | 1.97 |
| ATOM | 1555 | ŏ | | | 14.674 | | -13.372 | 1.00 | 0.28 |
| ATOM | | | SER | 106 | 14.669 | 1.781 | -14.006 | 1.00 | 0.31 |
| | 1556 | N | SER | 107 | 15.715 | 3.187 | -12.677 | 1.00 | 0.26 |
| MOTA | 1557 | HN | SER | 107 | 15.687 | | -12.166 | 1.00 | 0.25 |
| MOTA | 1558 | CA | SER | 107 | 16.940 | | -12.641 | | |
| MOTA | 1559 | HA | SER | 107 | | | | 1.00 | 0.27 |
| ATOM | 1560 | CB | | | 17.018 | | -13.560 | 1.00 | 0.29 |
| | | | SER | 107 | 18.175 | 3.226 | -12.474 | 1.00 | 0.28 |
| MOTA | 1561 | HB1 | SER | 107 | 18.292 | 3.847 | -13.353 | 1.00 | 1.12 |
| MOTA | 1562 | HB2 | SER | 107 | 19.049 | | -12.355 | | |
| ATOM | 1563 | OG | SER | 107 | | | | 1.00 | 1.04 |
| ATOM | 1564 | | | | 18.017 | | -11.320 | 1.00 | 1.29 |
| | | HG | SER | 107 | 18.556 | 4.827 | -11.436 | 1.00 | 1.82 |
| ATOM | 1565 | C | SER | 107 | 16.836 | | -11.460 | 1.00 | 0.26 |
| MOTA | 1566 | 0 | SER | 107 | 15.829 | | -10.781 | | |
| MOTA | 1567 | N | SER | 108 | | | | 1.00 | 0.26 |
| ATOM | 1568 | | | | 17.859 | | -11.203 | 1.00 | 0.28 |
| | | HN | SER | 108 | 18.666 | | -11.757 | 1.00 | 0.31 |
| MOTA | 1569 | CA | SER | 108 | 17.788 | | -10.061 | 1.00 | 0.30 |
| MOTA | 1570 | HA | SER | 108 | 16.775 | -0.706 | | | |
| MOTA | 1571 | CB | SER | 108 | | | | 1.00 | 0.30 |
| MOTA | 1572 | | | | 18.728 | | -10.330 | 1.00 | 0.36 |
| | | HB1 | SER | 108 | 19.561 | -1.505 | | 1.00 | 1.09 |
| ATOM | 1573 | | SER | 108 | 19.103 | | -11.338 | 1.00 | 0.95 |
| MOTA | 1574 | OG | SER | 108 | 18.005 | | -10.176 | | |
| ATOM | 1575 | HG | SER | 108 | 18.550 | | | 1.00 | 1.47 |
| ATOM | 1576 | c | | | | | -10.513 | 1.00 | 2.00 |
| ATOM | | | SER | 108 | 18.181 | 0.390 | -8.767 | 1.00 | 0.28 |
| | 1577 | 0 | SER | 108 | 19.279 | 0.265 | -8.261 | 1.00 | 0.33 |
| MOTA | 1578 | N | LYS | 109 | 17.272 | 1.157 | -8.224 | 1.00 | 0.24 |
| MOTA | 1579 | HN | LYS | 109 | 16 302 | 1 241 | 0 646 | 1.00 | 0.24 |
| | | | | | | | | | - |

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| ATOM | 1580 | CA | LYS | 109 | 17.561 | 1.897 | -6.960 | 1.00 | 0.23 |
|------|------|------|---------|------|--------|--------|---------|------|------|
| ATOM | 1581 | HA | LYS | 109 | 18.275 | 1.341 | -6.370 | | |
| | | | | | | | _ | 1.00 | 0.25 |
| MOTA | 1582 | CB | LYS | 109 | 18.123 | 3.293 | -7.268 | 1.00 | 0.24 |
| ATOM | 1583 | HB1 | LYS | 109 | 18.172 | 3.868 | -6.355 | 1.00 | 0.27 |
| MOTA | 1584 | HB2 | LYS | 109 | 17.472 | 3.793 | -7.970 | 1.00 | 0.25 |
| MOTA | 1585 | | LYS | | | | | | |
| | | | | 109 | 19.525 | 3.177 | -7.868 | 1.00 | 0.30 |
| MOTA | 1586 | HG1 | LYS | 109 | 19.476 | 2.615 | -8.785 | 1.00 | 0.54 |
| ATOM | 1587 | HG2 | LYS | 109 | 20.177 | 2.675 | -7.170 | 1.00 | 0.70 |
| ATOM | 1588 | | LYS | 109 | | | | | |
| | | | | | 20.072 | 4.574 | -8.169 | 1.00 | 0.75 |
| MOTA | 1589 | HD1 | LYS | 109 | 20.124 | 5.144 | -7.254 | 1.00 | 1.27 |
| ATOM | 1590 | HD2 | LYS | 109 | 19.420 | 5.074 | -8.870 | 1.00 | 1.27 |
| ATOM | 1591 | CE | LYS | 109 | 21.475 | 4.453 | | | |
| | | | | | | | -8.770 | 1.00 | 1.13 |
| MOTA | 1592 | HE1 | | 109 | 21.396 | 4.264 | -9.830 | 1.00 | 1.68 |
| MOTA | 1593 | HE2 | LYS | 109 | 22.000 | 3.636 | -8.297 | 1.00 | 1.68 |
| MOTA | 1594 | NZ | LYS | 109 | 22.224 | 5.721 | -8.545 | 1.00 | 1.79 |
| MOTA | 1595 | HZ1 | _ | 109 | 21.689 | | | | |
| | | | | | | 6.516 | -8.948 | 1.00 | 2.22 |
| ATOM | 1596 | HZ2 | | 109 | 23.155 | 5.660 | -9.006 | 1.00 | 2.17 |
| ATOM | 1597 | HZ3 | LYS | 109 | 22.351 | 5.873 | -7.525 | 1.00 | 2.34 |
| ATOM | 1598 | C | LYS | 109 | 16.259 | 2.052 | -6.175 | 1.00 | 0.21 |
| ATOM | 1599 | Ō | LYS | 109 | 15.190 | | | | |
| | | | | | | 2.110 | -6.747 | 1.00 | 0.20 |
| ATOM | 1600 | N | GLY | 110 | 16.338 | 2.124 | -4.873 | 1.00 | 0.23 |
| MOTA | 1601 | HN | GLY | 110 | 17.212 | 2.079 | -4.432 | 1.00 | 0.26 |
| MOTA | 1602 | CA | GLY | 110 | 15.099 | 2.283 | -4.056 | 1.00 | |
| MOTA | 1603 | HA1 | GLY | 110 | | | | | 0.22 |
| | | | | | 14.751 | 3.302 | -4.124 | 1.00 | 0.23 |
| MOTA | 1604 | HA2 | GLY | 110 | 15.316 | 2.044 | -3.024 | 1.00 | 0.25 |
| MOTA | 1605 | C | GLY | 110 | 14.013 | 1.342 | -4.581 | 1.00 | 0.19 |
| ATOM | 1606 | 0 | GLY | 110 | 14.281 | | | | |
| | | | | | | 0.216 | -4.949 | 1.00 | 0.20 |
| MOTA | 1607 | N | TYR | 111 | 12.789 | 1.801 | -4.626 | 1.00 | 0.17 |
| MOTA | 1608 | HN | TYR | 111 | 12.599 | 2.716 | -4.330 | 1.00 | 0.18 |
| MOTA | 1609 | CA | TYR | 111 | 11.683 | 0.941 | -5.136 | 1.00 | |
| ATOM | 1610 | HA | TYR | 111 | | | | | 0.15 |
| | | | | | 11.975 | -0.098 | -5.088 | 1.00 | 0.16 |
| MOTA | 1611 | CB | TYR | 111 | 10.437 | 1.162 | -4.277 | 1.00 | 0.15 |
| MOTA | 1612 | HB1 | TYR | 111 | 9, 633 | 0.540 | -4.641 | 1.00 | 0.15 |
| MOTA | 1613 | HB2 | TYR | 111 | 10.143 | 2.200 | | | |
| MOTA | 1614 | | | | | | -4.330 | 1.00 | 0.16 |
| | | CG | TYR | 111 | 10.745 | 0.798 | -2.844 | 1.00 | 0.17 |
| ATOM | 1615 | CD1 | TYR | 111 | 10.648 | -0.533 | -2.422 | 1.00 | 0.17 |
| MOTA | 1616 | HD1 | TYR | 111 | 10.354 | -1.301 | -3.121 | 1.00 | 0.17 |
| ATOM | 1617 | | TYR | 111. | | | | | |
| | | | | | 11.127 | 1.794 | -1.936 | 1.00 | 0.20 |
| MOTA | 1618 | | TYR | 111 | 11.201 | 2.821 | -2.261 | 1.00 | 0.23 |
| MOTA | 1619 | CE1 | TYR | 111 | 10.933 | -0.868 | -1.093 | 1.00 | 0.19 |
| MOTA | 1620 | HE1 | TYR | 111 | 10.858 | -1.895 | -0.767 | | |
| MOTA | 1621 | | | | | - | | 1.00 | 0.20 |
| | | CE2 | | 111 | 11.412 | 1.459 | -0.607 | 1.00 | 0.22 |
| MOTA | 1622 | HE2 | TYR | 111 | 11.706 | 2.227 | 0.093 | 1.00 | 0.26 |
| ATOM | 1623 | CZ | TYR | 111 | 11.315 | 0.127 | -0.185 | 1.00 | |
| MOTA | 1624 | OH | TYR | 111 | | | | | 0.21 |
| | | | | | 11.595 | -0.204 | 1.125 | 1.00 | 0.23 |
| MOTA | 1625 | HH | TYR | 111 | 12.543 | -0.121 | 1.255 | 1.00 | 0.95 |
| ATOM | 1626 | C | TYR | 111 | 11.374 | 1.321 | -6.588 | 1.00 | 0.14 |
| ATOM | 1627 | 0 | TYR | 111 | 10.949 | 2.424 | -6.871 | 1.00 | |
| MOTA | 1628 | N | ASN | | | | | | 0.15 |
| | | | | 112 | 11.581 | 0.421 | -7.511 | 1.00 | 0.15 |
| MOTA | 1629 | HN | ASN | 112 | 11.924 | -0.464 | -7.264 | 1.00 | 0.17 |
| MOTA | 1630 | CA | ASN | 112 | 11.295 | 0.739 | -8.939 | 1.00 | 0.16 |
| ATOM | 1631 | HA | ASN | 112 | 11.870 | 1.605 | -9.235 | | |
| MOTA | 1632 | CB | ASN | | | | | 1.00 | 0.16 |
| | | | | 112 | 11.677 | -0.450 | -9.822 | 1.00 | 0.19 |
| ATOM | 1633 | HB1 | | 112 | 11.025 | -1.276 | -9.607 | 1.00 | 0.22 |
| ATOM | 1634 | HB2 | asn | 112 | 12.698 | -0.739 | -9.622 | 1.00 | 0.19 |
| MOTA | 1635 | CG | ASN | 112 | 11.531 | | -11.295 | | |
| MOTA | 1636 | OD1 | ASN | 112 | | | | 1.00 | 0.24 |
| | | | | | 10.446 | | -11.748 | 1.00 | 0.96 |
| MOTA | 1637 | | ASN | 112 | 12.583 | ~0.059 | -12.067 | 1.00 | 1.06 |
| MOTA | 1638 | HD21 | ASN | 112 | 13.458 | -0.308 | -11.704 | 1.00 | 1.80 |
| MOTA | 1639 | HD22 | ASN | 112 | 12.497 | | -13.012 | 1.00 | |
| MOTA | 1640 | C | ASN | | | | | | 1.08 |
| | | | | 112 | 9.803 | 1.040 | | 1.00 | 0.15 |
| MOTA | 1641 | 0 | asn | 112 | 8.953 | 0.310 | -8.637 | 1.00 | 0.14 |
| ATOM | 1642 | N | LEU | 113 | 9.482 | | -9.777 | 1.00 | 0.15 |
| MOTA | 1643 | HN | LEU | 113 | 10.187 | | | | |
| ATOM | | | | | | 2.684 | | 1.00 | 0.16 |
| | 1644 | CA | LEU | 113 | 8.049 | 2.475 | | 1.00 | 0.15 |
| ATOM | 1645 | HA | LEU | 113 | 7.582 | 2.620 | -9.025 | 1.00 | 0.14 |
| MOTA | 1646 | CB | LEU | 113 | 7.981 | | -10.791 | 1.00 | 0.16 |
| MOTA | 1647 | HB1 | | 113 | | | | | |
| | | | | | 8.513 | | -11.721 | 1.00 | 0.17 |
| MOTA | 1648 | | LEU | 113 | 8.452 | 4.571 | -10.226 | 1.00 | 0.16 |
| MOTA | 1649 | CG | LEU | 113 | 6.523 | | -11.095 | 1.00 | 0.17 |
| ATOM | 1650 | HG | LEU | 113 | 6.041 | | | | |
| ATOM | 1651 | | | | | | -11.652 | 1.00 | 0.18 |
| | | | LEU | 113 | 5.748 | 4.421 | | 1.00 | 0.18 |
| MOTA | | HD11 | - | 113 | 4.841 | 4.969 | -10.007 | 1.00 | 0.99 |
| MOTA | 1653 | HD12 | LEU | 113 | 6.359 | 4.991 | | 1.00 | 1.00 |
| MOTA | 1654 | HD13 | | 113 | 5.490 | | | | |
| ATOM | | | | | | 3.474 | | 1.00 | 0.97 |
| | 1655 | | LEU | 113 | 6.526 | | -11.943 | 1.00 | 0.20 |
| MOTA | 1656 | HD21 | र एच. र | 113 | £ 115 | E 277 | _11 274 | 1 00 | 1 05 |

| ATOM | | HD22 | | 113 | 5.930 | 5.302 -12.83 | 1.00 | 1.03 |
|--------------|--------------|------------|------------|-----|--------|-----------------|---------|------|
| MOTA | 1658 | HD23 | LEU | 113 | 7.539 | 5.696 -12.233 | 1.00 | 1.00 |
| MOTA | 1659 | С | LEU | 113 | 7.320 | 1.361 -10.743 | | 0.15 |
| MOTA | 1660 | 0 | LEU | 113 | 6.203 | 1.014 -10.419 | | 0.15 |
| MOTA | 1661 | N | PHE | 114 | 7.928 | 0.817 -11.762 | | 0.16 |
| MOTA | 1662 | HN | PHE | 114 | 8.822 | 1.123 -12.020 | | 0.17 |
| MOTA | 1663 | CA | PHE | 114 | 7.245 | -0.250 -12.55 | | 0.17 |
| MOTA | 1664 | HA | PHE | 114 | 6.338 | 0.151 -12.980 | | 0.18 |
| MOTA | 1665 | CB | PHE | 114 | 8.159 | -0.720 -13.685 | | 0.21 |
| MOTA | 1666 | HB1 | PHE | 114 | 9.077 | -1.108 -13.27 | | 0.22 |
| MOTA | 1667 | HB2 | PHE | 114 | 8.380 | 0.111 -14.340 | | 0.22 |
| MOTA | 1668 | CG | PHE | 114 | 7.457 | -1.807 -14.46 | | 0.24 |
| MOTA | 1669 | CD1 | PHE | 114 | 7.545 | -3.135 -14.03 | | 0.35 |
| MOTA | 1670 | HD1 | PHE | 114 | 8.105 | -3.376 -13.14 | | 0.43 |
| MOTA | 1671 | CD2 | PHE | 114 | 6.724 | -1.494 -15.61 | | 0.24 |
| MOTA | 1672 | HD2 | PHE | 114 | 6.655 | -0.470 -15.950 | | 0.28 |
| MOTA | 1673 | CE1 | PHE | 114 | 6.902 | -4.149 -14.74 | _ + + + | 0.39 |
| MOTA | 1674 | HE1 | PHE | 114 | 6.975 | -5.171 -14.402 | | 0.50 |
| MOTA | 1675 | CE2 | PHE | 114 | 6.078 | -2.512 -16.32 | | 0.26 |
| MOTA | 1676 | HE2 | PHE | 114 | 5.511 | -2.273 -17.214 | | 0.30 |
| MOTA | 1677 | CZ | PHE | 114 | 6.168 | -3.839 -15.890 | | 0.32 |
| MOTA | 1678 | HZ | PHE | 114 | 5.670 | -4.623 -16.438 | | 0.35 |
| MOTA | 1679 | C | PHE | 114 | 6.900 | -1.452 -11.67 | | 0.17 |
| MOTA | 1680 | 0 | PHE | 114 | 5.842 | -2.034 -11.80 | | 0.17 |
| MOTA | 1681 | N | LEU | 115 | 7.774 | -1.846 -10.79 | | 0.18 |
| MOTA | 1682 | HN | LEU | 115 | 8.631 | -1.380 -10.70 | | 0.18 |
| ATOM | 1683 | CA | LEU | 115 | 7.463 | -3.028 -9.94 | | 0.20 |
| MOTA | 1684 | HA | LEU | 115 | 7.297 | -3.882 -10.579 | | 0.21 |
| MOTA | 1685 | CB | LEU | 115 | 8.634 | -3.304 -8.984 | | 0.23 |
| ATOM | 1686 | HB1 | LEU | 115 | 8.237 | -3.650 -8.04: | | 0.26 |
| MOTA | 1687 | HB2 | LEU | 115 | 9.172 | -2.387 -8.82 | | 0.22 |
| MOTA | 1688 | CG | LEU | 115 | 9.612 | -4.369 -9.539 | | 0.28 |
| MOTA | 1689 | HG | LEU | 115 | 10.397 | -4.525 -8.812 | | 0.33 |
| ATOM | 1690 | | LEU | 115 | 8.886 | -5.702 -9.749 | | 0.36 |
| MOTA | 1691 | | _ | 115 | 9.551 | -6.514 -9.498 | | 0.99 |
| ATOM | | | _ | 115 | 8.578 | -5.795 -10.779 | | 1.11 |
| ATOM | 1693 | HD13 | - | 115 | 8.017 | -5.740 -9.10 | | 1.13 |
| MOTA | 1694 | | LEU | 115 | 10.249 | -3.903 -10.85 | | 0.30 |
| MOTA | | HD21 | | 115 | 10.497 | -4.761 -11.46 | | 1.10 |
| ATOM | 1696 | HD22 | | 115 | 11.149 | -3.351 -10.649 | | 1.06 |
| MOTA | 1697 | HD23 | LEU | 115 | 9.567 | -3.272 -11.39 | | 1.01 |
| MOTA | 1698 | C | LEU | 115 | 6.194 | -2.748 - 9.136 | | 0.19 |
| MOTA | 1699 | 0 | LEU | 115 | 5.280 | -3.548 -9.10 | | 0.20 |
| ATOM | 1700 | N | VAL | 116 | 6.130 | -1.624 -8.47 | | 0.18 |
| ATOM | 1701 | HN | VAL | 116 | 6.879 | -0.993 -8.508 | • | 0.18 |
| MOTA | 1702 | CA | VAL | 116 | 4.919 | -1.305 -7.664 | | 0.19 |
| ATOM | 1703 | HA | VAL | 116 | 4.686 | -2.146 -7.028 | | 0.21 |
| ATOM | 1704 | CB | VAL | 116 | 5.203 | -0.078 -6.794 | | 0.20 |
| ATOM | 1705 | HB | VAL | 116 | 5.581 | 0.722 -7.414 | | 0.19 |
| MOTA | 1706 | | VAL | 116 | 3.914 | 0.381 - 6.103 | | 0.22 |
| MOTA | 1707 | HG11 | | 116 | 3.253 | 0.832 -6.828 | | 1.05 |
| MOTA | | HG12 | | 116 | 4.155 | 1.105 -5.339 | | 1.05 |
| MOTA | | HG13 | | 116 | 3.426 | -0.470 -5.650 | | 1.03 |
| MOTA | 1710 | | VAL | 116 | 6.246 | -0.443 -5.73 | | 0.21 |
| MOTA | | HG21 | | 116 | 7.188 | -0.654 -6.223 | | 1.02 |
| MOTA | | HG22 | | 116 | 5.917 | -1.317 -5.194 | | 0.98 |
| ATOM | 1713 | HG23 | | 116 | 6.370 | 0.382 -5.052 | | 1.03 |
| MOTA | 1714 | C | VAL | 116 | 3.724 | -1.020 -8.582 | | 0.18 |
| MOTA | 1715 | 0 | VAL | 116 | 2.615 | -1.433 - 8.312 | | 0.19 |
| MOTA | 1716 | N | ALA | 117 | 3.934 | -0.307 - 9.659 | | 0.17 |
| MOTA | 1717 | HN | ALA | 117 | 4.833 | 0.028 -9.859 | | 0.16 |
| MOTA | 1718 | CA | ALA | 117 | 2.796 | 0.007 -10.572 | 2 1.00 | 0.17 |
| ATOM | 1719 | HA | ALA | 117 | 2.064 | 0.598 - 10.044 | 1.00 | 0.19 |
| MOTA MOTA | 1720 | CB | ALA | 117 | 3.306 | 0.795 -11.780 | 1.00 | 0.18 |
| ATOM | 1721 | | ALA | 117 | 4.378 | 0.709 - 11.840 | | 1.05 |
| | 1722 | | ALA | 117 | 3.033 | 1.834 -11.674 | 1.00 | 1.01 |
| MOTA MOTA | 1723 | HB3 | ALA | 117 | 2.863 | 0.397 -12.682 | 1.00 | 0.98 |
| ATOM | 1724 | C | ALA | 117 | 2.150 | -1.291 -11.058 | 1.00 | 0.17 |
| ATOM | 1725 | 0 | ALA | 117 | 0.956 | -1.480 -10.951 | 1.00 | 0.19 |
| ATOM | 1726 | N | ALA | 118 | 2.931 | -2.187 - 11.588 | 1.00 | 0.16 |
| ATOM | 1727 | HN | ALA | 118 | 3.893 | -2.015 - 11.663 | 1.00 | 0.16 |
| ATOM | 1728 | CA | ALA | 118 | 2.366 | -3.472 - 12.083 | 1.00 | 0.17 |
| MOTA | 1729 1730 | HA | ALA | 118 | 1.643 | -3.273 -12.859 | 1.00 | 0.19 |
| ATOM | 1731 | CB | ALA | 118 | 3.491 | -4.335 -12.653 | 1.00 | 0.17 |
| ATOM | 1732 | HB1 HB2 | | 118 | 3.125 | -5.338 -12.812 | | 1.05 |
| ATOM | 1733 | HB3 | | 118 | 4.316 | -4.358 -11.956 | | 1.02 |
| 404 | _, | ***** | STIFF. | 118 | 3 824 | -1 03U -13 E03 | 1 00 | 1 07 |

| ATOM | 1734 | C A | LA 1 | 118 | 1.687 | -4.220 | -10.935 | 1.00 | 0.17 |
|------|--------|--------|------------|-----|--------|--------------------|---------|------|------|
| MOTA | 1735 | O A | LA 1 | 18 | 0.699 | | -11.124 | 1.00 | 0.18 |
| MOTA | 1736 | | | 19 | | _ | | | |
| | | _ | | | 2.225 | -4.123 | -9.751 | 1.00 | 0.16 |
| MOTA | 1737 | | | 119 | 3.035 | -3.585 | -9.623 | 1.00 | 0.16 |
| MOTA | 1738 | CA H | IS 1 | 19 | 1.627 | -4.855 | -8.599 | 1.00 | 0.17 |
| MOTA | 1739 | HA H | IS 1 | 119 | 1.576 | -5.907 | -8.833 | 1.00 | 0.18 |
| MOTA | 1740 | | | 119 | 2.513 | -4.655 | | | |
| | | | | | | | -7.368 | 1.00 | 0.19 |
| MOTA | 1741 | | | 19 | 2.547 | -3.605 | -7.116 | 1.00 | 0.19 |
| MOTA | 1742 | нв2 н | IS 1 | L19 | 3.512 | -5.005 | -7.584 | 1.00 | 0.20 |
| MOTA | 1743 | CG H | IS 1 | L19 | 1.950 | -5.431 | -6.210 | 1.00 | 0.21 |
| ATOM | 1744 | ND1 H | | 119 | | | | | |
| | | | | | 2.228 | -6.775 | -6.020 | 1.00 | 0.26 |
| MOTA | 1745 | HD1 .H | | 119 | 2.791 | -7.336 | -6.593 | 1.00 | 0.30 |
| MOTA | 1746 | CD2 H | IIS 3 | 119 | 1.128 | -5.067 | -5.172 | 1.00 | 0.20 |
| MOTA | 1747 | HD2 H | IIS 1 | 119 | 0.719 | -4.079 | -5.019 | 1.00 | 0.21 |
| MOTA | 1748 | | | 119 | | -7.168 | | | |
| | | | | | 1.585 | | -4.906 | 1.00 | 0.27 |
| ATOM | 1749 | HE1 H | | 119 | 1.622 | -8.171 | -4.509 | 1.00 | 0.33 |
| MOTA | 1750 | NE2 H | IIS I | 119 | 0.899 | -6.166 | -4.350 | 1.00 | 0.23 |
| ATOM | 1751 | C H | IIS 3 | 119 | 0.215 | -4.333 | -8.299 | 1.00 | 0.17 |
| MOTA | 1752 | | | 119 | | | | | |
| | | | | | -0.721 | -5.101 | -8.185 | 1.00 | 0.18 |
| ATOM | 1753 | | | 120 | 0.043 | -3.044 | -8.160 | 1.00 | 0.18 |
| MOTA | 1754 | HN G | SLU : | 120 | 0.801 | -2.430 | -8.248 | 1.00 | 0.18 |
| MOTA | 1755 | CA G | LU : | 120 | -1.322 | -2.520 | -7.860 | 1.00 | 0.20 |
| MOTA | 1756 | | | 120 | -1.666 | -2.977 | | | |
| | | | | | | | -6.943 | 1.00 | 0.21 |
| MOTA | 1757 | | | 120 | -1.294 | -0.999 | -7.668 | 1.00 | 0.22 |
| MOTA | 1758 | HB1 G | LU : | 120 | -0.719 | -0.763 | -6.785 | 1.00 | 0.37 |
| MOTA | 1759 | HB2 G | LU : | 120 | -2.302 | -0.635 | -7.542 | 1.00 | 0.33 |
| MOTA | 1760 | | | 120 | -0.663 | -0.314 | | | |
| ATOM | | | | | | | -8.875 | 1.00 | 0.41 |
| | 1761 | | | 120 | -1.125 | -0.668 | -9.781 | 1.00 | 0.63 |
| MOTA | 1762 | HG2 G | SLU : | 120 | 0.393 | -0.531 | -8.895 | 1.00 | 0.87 |
| ATOM | 1763 | CD G | SLU : | 120 | -0.875 | 1.194 | -8.757 | 1.00 | 0.94 |
| MOTA | 1764 | | | 120 | | | | | |
| | | _ | | | -0.757 | 1.703 | -7.654 | 1.00 | 1.67 |
| MOTA | 1765 | | | 120 | -1.151 | 1.816 | -9.769 | 1.00 | 1.56 |
| MOTA | 1766 | C | SLU : | 120 | -2.291 | -2.903 | -8.984 | 1.00 | 0.20 |
| ATOM | 1767 | 0 0 | GLU : | 120 | -3.432 | -3.238 | | 1.00 | 0.21 |
| MOTA | 1768 | | | 121 | | | | | |
| | | | | | -1.853 | | -10.217 | 1.00 | 0.19 |
| ATOM | 1769 | • | | 121 | -0.928 | -2.608 | -10.405 | 1.00 | 0.19 |
| MOTA | 1770 | CA E | PHE | 121 | -2.767 | -3.251 | -11.331 | 1.00 | 0.21 |
| MOTA | 1771 | HA I | PHE | 121 | -3.628 | | -11.317 | 1.00 | 0.23 |
| MOTA | 1772 | | | 121 | | | | | |
| | | | | | -2.053 | | -12.685 | 1.00 | 0.22 |
| | . 1773 | | | 121 | -2.576 | -3.726 | -13.419 | 1.00 | 0.24 |
| ATOM | 1774 | HB2 I | PHE | 121 | -1.041 | -3.493 | -12.587 | 1.00 | 0.21 |
| MOTA | 1775 | CG I | PHE | 121 | -2.026 | | -13.141 | 1.00 | 0.25 |
| ATOM | 1776 | | | 121 | | | | | |
| | | _ | | | -0.804 | | -13.308 | 1.00 | 0.27 |
| ATOM | 1777 | | | 121 | 0.121 | -1.535 | -13.113 | 1.00 | 0.40 |
| MOTA | 1778 | CD2 I | PHE | 121 | -3.227 | -1.007 | -13.403 | 1.00 | 0.45 |
| MOTA | 1779 | HD2 I | PHE | 121 | -4.173 | | -13.281 | 1.00 | 0.60 |
| MOTA | 1780 | | | 121 | | | | | |
| | | | | | -0.781 | | -13.733 | 1.00 | 0.29 |
| MOTA | 1781 | | | 121 | 0.163 | 0.824 | -13.862 | 1.00 | 0.39 |
| MOTA | 1782 | CE2 I | PHE | 121 | -3.202 | 0.327 | -13.828 | 1.00 | 0.49 |
| MOTA | 1783 | HE2 I | PHE | 121 | -4.127 | 0.847 | -14.029 | 1.00 | 0.68 |
| MOTA | 1784 | CZ I | | 121 | -1.979 | | -13.993 | | |
| ATOM | 1785 | | | | | | | 1.00 | 0.34 |
| | | | | 121 | -1.961 | | -14.321 | 1.00 | 0.38 |
| MOTA | 1786 | C | | 121 | -3.228 | -4.693 | -11.120 | 1.00 | 0.20 |
| MOTA | 1787 | 0 1 | PHE | 121 | -4.374 | -5.027 | -11.344 | 1.00 | 0.21 |
| ATOM | 1788 | N C | GLY | 122 | -2.344 | | -10.690 | 1.00 | 0.18 |
| MOTA | 1789 | | | 122 | -1.424 | | | | |
| ATOM | 1790 | | | | | | -10.514 | 1.00 | 0.17 |
| | | | | 122 | -2.737 | | -10.464 | 1.00 | 0.20 |
| MOTA | 1791 | HA1 (| GLY | 122 | -1.890 | -7.523 | -10.092 | 1.00 | 0.21 |
| ATOM | 1792 | HA2 (| GLY | 122 | -3.072 | -7.404 | -11.394 | 1.00 | 0.21 |
| ATOM | 1793 | | | 122 | -3.867 | -7.022 | _ | | |
| MOTA | 1794 | | | | | | | 1.00 | 0.20 |
| | | | | 122 | -4.823 | - 7.756 | -9.589 | 1.00 | 0.22 |
| MOTA | 1795 | N i | HIS | 123 | -3.778 | -6.240 | -8.392 | 1.00 | 0.20 |
| MOTA | 1796 | HN F | HIS | 123 | -3.005 | -5.644 | -8.287 | 1.00 | 0.20 |
| MOTA | 1797 | | | 123 | -4.864 | | | | |
| ATOM | | | | | _ | -6.243 | -7.371 | 1.00 | 0.22 |
| | 1798 | | | 123 | -5.047 | -7.255 | -7.042 | 1.00 | 0.23 |
| MOTA | 1799 | | | 123 | -4.456 | -5.382 | -6.174 | 1.00 | 0.25 |
| ATOM | 1800 | HB1 F | HIS | 123 | -5.324 | -5.180 | -5.564 | 1.00 | 0.30 |
| MOTA | 1801 | _ | | 123 | -4.041 | -4.449 | _ | | |
| MOTA | 1802 | | | | | | -6.527 | 1.00 | 0.25 |
| | | | | 123 | -3.427 | -6.108 | -5.354 | 1.00 | 0.27 |
| MOTA | 1803 | ND1 I | | 123 | -3.736 | -7.247 | -4.628 | 1.00 | 0.37 |
| MOTA | 1804 | HD1 F | HIS | 123 | -4.611 | -7.685 | -4.581 | 1.00 | 0.45 |
| ATOM | 1805 | CD2 I | | 123 | -2.096 | -5.866 | -5.125 | | |
| MOTA | 1806 | HD2 I | | | | | | 1.00 | 0.25 |
| ATOM | | | | 123 | -1.532 | -5.046 | -5.545 | 1.00 | 0.27 |
| | 1807 | CE1 E | | 123 | -2.614 | -7.644 | -4.001 | 1.00 | 0.38 |
| MOTA | 1808 | HE1 F | | 123 | ~2.553 | -8.514 | -3.367 | 1.00 | 0.47 |
| MOTA | 1809 | NE2 1 | HIS | 123 | -1.584 | -6.837 | -4.269 | 1.00 | 0.29 |
| MOTA | 1810 | | | 123 | -6 137 | -5 671 | -7 007 | 1 00 | 0.23 |
| | | - | _ | | | | | | |

| • | | | | | | | | | |
|------|------|------|------------|-------|--------|---------|---------|------|------|
| MOTA | 1811 | 0 | HIS | 123 | -7.229 | -6.148 | -7.755 | 1.00 | 0.25 |
| MOTA | 1812 | _ | SER | 124 | -6.002 | -4.646 | -8.788 | 1.00 | |
| | _ | | | | | • • | | | 0.23 |
| MOTA | 1813 | | SER | 124 | -5.110 | -4.278 | -8.962 | 1.00 | 0.22 |
| MOTA | 1814 | CA | SER | 124 | -7.196 | -4.030 | -9.429 | 1.00 | 0.25 |
| MOTA | 1815 | HA | SER | 124 | -7.928 | -3.790 | -8.672 | 1.00 | 0.27 |
| MOTA | 1816 | | SER | 124 | | | - | | |
| | | | | | -6.778 | | -10.156 | 1.00 | 0.27 |
| ATOM | 1817 | HB1 | SER | 124 | -6.219 | -2.119 | -9.478 | 1.00 | 0.29 |
| ATOM | 1818 | HB2 | SER | 124 | -7.654 | -2.224 | -10.494 | 1.00 | 0.29 |
| MOTA | 1819 | | SER | 124 | -5.975 | | -11.279 | 1.00 | 0.25 |
| | | | | | | | | | |
| MOTA | 1820 | HG | SER | 124 | -6.545 | -3.131 | -12.050 | 1.00 | 0.88 |
| MOTA | 1821 | C. | SER | 124 | -7.805 | -5.006 | -10.437 | 1.00 | 0.24 |
| ATOM | 1822 | | SER | 124 | -8.975 | | -10.755 | 1.00 | 0.26 |
| | | | | | | | | | |
| MOTA | 1823 | | LEU | 125 | -7.022 | | -10.952 | 1.00 | 0.22 |
| MOTA | 1824 | HN | LEU | 125 | -6.078 | -5.953 | -10.690 | 1.00 | 0.21 |
| MOTA | 1825 | CA | LEU | 125 | -7.562 | | -11.949 | 1.00 | 0.23 |
| ATOM | 1826 | | | | | | | | |
| | | | LEU | 125 | -8.285 | | -12.568 | 1.00 | 0.24 |
| MOTA | 1827 | CB | LEU | 125 | -6.420 | -7.398 | -12.827 | 1.00 | 0.22 |
| MOTA | 1828 | HB1 | LEU | 125 | -6.759 | -8.247 | -13.398 | 1.00 | 0.24 |
| MOTA | 1829 | HB2 | | 125 | -5.594 | | -12.197 | | |
| | | | | | | _ | | 1.00 | 0.22 |
| ATOM | 1830 | CG | LEU | 125 | -5.956 | -6.280 | -13.779 | 1.00 | 0.22 |
| MOTA | 1831 | HG | LEŲ | 125 | -5.928 | -5.343 | -13.241 | 1.00 | 0.24 |
| MOTA | 1832 | CD1 | | 125 | -4.556 | | -14.302 | 1.00 | 0.25 |
| | | | | | | | | | |
| MOTA | 1833 | HD11 | - | 125 | -4.588 | | -14.874 | 1.00 | 0.99 |
| MOTA | | HD12 | LEU | 125 | -3.879 | -6.719 | -13.471 | 1.00 | 1.00 |
| MOTA | 1835 | HD13 | LEU | · 125 | -4.215 | -5.794 | -14.933 | 1.00 | 1.05 |
| MOTA | 1836 | CD2 | | 125 | -6.913 | | -14.976 | | |
| | | | | | | | | 1.00 | 0.24 |
| MOTA | | HD21 | | 125 | -7.793 | -5.604 | -14.682 | 1.00 | 1.05 |
| MOTA | 1838 | HD22 | LEU | 125 | -7.201 | -7.135 | -15.324 | 1.00 | 1.00 |
| ATOM | | | | 125 | -6.415 | | -15.775 | 1.00 | 1.03 |
| | | | | | | | | | |
| MOTA | 1840 | C | LEU | 125 | -8.256 | -8.044 | -11.234 | 1.00 | 0.24 |
| MOTA | 1841 | 0 | LEU | 125 | -8.790 | -8.935 | -11.864 | 1.00 | 0.33 |
| ATOM | 1842 | N | GLY | 126 | -8.277 | -8.035 | -9.927 | 1.00 | 0.24 |
| MOTA | 1843 | HN | | 126 | | | | | |
| | | | GLY | | -7.858 | -7.298 | | 1.00 | 0.29 |
| MOTA | 1844 | CA | GLY | 126 | -8.968 | -9.132 | -9.185 | 1.00 | 0.27 |
| MOTA | 1845 | HA1 | GLY | 126 | -9.748 | -9.545 | -9.807 | 1.00 | 0.29 |
| ATOM | 1846 | _ | GLY | 126 | -9.408 | -8.727 | | | |
| | | | | | | | | 1.00 | 0.29 |
| MOTA | 1847 | С | GLY | 126 | -7.985 | -10.245 | -8.809 | 1.00 | 0.26 |
| MOTA | 1848 | 0 | GLY | 126 | -8.377 | -11,268 | -8.283 | 1.00 | 0.30 |
| MOTA | 1849 | N | LEU | 127 | | -10.068 | | 1.00 | 0.23 |
| | | | | | | | | | |
| MOTA | 1850 | HN | LEU | 127 | | -9.239 | | 1.00 | 0.22 |
| MOTA | 1851 | CA | LEU | 127 | -5.744 | -11.138 | -8.700 | 1.00 | 0.25 |
| MOTA | 1852 | HA | LEU | 127 | | -12.099 | | 1.00 | 0.28 |
| ATOM | | | | | | | | | |
| _ | 1853 | CB | LEU | 127 | _ | -11.052 | | 1.00 | 0.23 |
| MOTA | 1854 | HB1 | LEU | 127 | -3.733 | -11.696 | -9.211 | 1.00 | 0.25 |
| MOTA | 1855 | HB2 | LEU | 127 | -4.156 | -10.033 | -9.602 | 1.00 | 0.22 |
| MOTA | 1856 | CG | LEU | 127 | | | | | |
| | | | | | | | -11.045 | 1.00 | 0.24 |
| MOTA | 1857 | HG | LEU | 127 | -5.707 | -10.915 | -11.384 | 1.00 | 0.23 |
| MOTA | 1858 | CD1 | LEU | 127 | -3.646 | -11,159 | -11.962 | 1.00 | 0.24 |
| MOTA | 1859 | HD11 | T.FIT | 127 | | | -12.868 | | |
| | | | | | | | | 1.00 | 1.00 |
| ATOM | 1860 | HD12 | | 127 | | | -12.208 | 1.00 | 1.02 |
| MOTA | 1861 | HD13 | LEU | . 127 | -2.962 | -10.491 | -11.460 | 1.00 | 1.03 |
| ATOM | 1862 | CD2 | LEU | 127 | | | -11.109 | 1.00 | 0.30 |
| MOTA | 1863 | HD21 | | 127 | | | | | |
| | | | | | | | -12.121 | 1.00 | 1.04 |
| ATOM | 1864 | HD22 | | 127 | -6.169 | -13.159 | -10.805 | 1.00 | 1.11 |
| MOTA | 1865 | HD23 | LEU | 127 | -4.478 | -13.515 | -10.454 | 1.00 | 1.03 |
| MOTA | 1866 | C | LEU | 127 | | • | -7.241 | 1.00 | 0.28 |
| ATOM | 1867 | | LEU | | | | | | |
| | | • | | 127 | | | -6.723 | | 0.32 |
| MOTA | 1868 | N | ASP | 128 | -5.027 | -12.059 | -6.581 | 1.00 | 0.32 |
| MOTA | 1869 | HN | ASP | 128 | -5.093 | -12.928 | -7.029 | 1.00 | 0.34 |
| MOTA | 1870 | CA | ASP | 128 | | -11.997 | · – | | |
| | | | | | | | - | 1.00 | 0.39 |
| MOTA | 1871 | HA | ASP | 128 | -4.882 | -11.046 | -4.728 | 1.00 | 0.40 |
| MOTA | 1872 | CB | ASP | 128 | -5.271 | -13.130 | -4.375 | 1.00 | 0.48 |
| MOTA | 1873 | HR1 | ASP | 128 | | -14.064 | • | | |
| ATOM | | | | | | | - | 1.00 | 0.48 |
| | 1874 | | ASP | 128 | | -13.193 | _ | 1.00 | 0.50 |
| MOTA | 1875 | CG | ASP | 128 | -5.171 | -12.854 | -2.873 | 1.00 | 0.55 |
| MOTA | 1876 | OD1 | ASP | 128 | | -12.980 | | | 1.23 |
| MOTA | 1877 | | ASP | 128 | | | • | | |
| | | | | | | -12.521 | _ | 1.00 | 1.22 |
| MOTA | 1878 | C | ASP | 128 | -3.078 | -12.159 | -5.082 | 1.00 | 0.37 |
| MOTA | 1879 | 0 | ASP | 128 | -2.424 | -12.387 | | 1.00 | 0.59 |
| ATOM | 1880 | N | HIS | 129 | | -12.042 | | | |
| | | | | | | _ | | 1.00 | 0.23 |
| MOTA | 1881 | HN | HIS | 129 | | -11.856 | | 1.00 | 0.32 |
| MOTA | 1882 | CA | HIS | 129 | -1.029 | -12.189 | -3.797 | 1.00 | 0.22 |
| MOTA | 1883 | HA | HIS | 129 | | -11.439 | • | 1.00 | 0.21 |
| MOTA | 1884 | | | | | | | | |
| | | CB | HIS | 129 | | -12.019 | | 1.00 | 0.23 |
| MOTA | 1885 | HB1 | HIS | 129 | | -12.302 | | 1.00 | 0.24 |
| MOTA | 1886 | HB2 | HIS | 129 | -1.217 | -12.653 | | 1.00 | 0.25 |
| ATOM | 1887 | CG | HIS | 129 | | -10.585 | | 1.00 | |
| | • | | | | V.113 | TO. 707 | 4.716 | 1.00 | 0.22 |

| | 1000 | | | | | | |
|--------------|--------------|--------------------|--------------|----------------------------------|------------------|--------------|--------------|
| MOTA | 1888 | ND1 HIS | 129 | -1.862 -10.161 | -1.156 | 1.00 | 0.35 |
| MOTA | 1889 | HD1 HIS | 129 | -2.602 -10.720 | -0.841 | 1.00 | 0.53 |
| MOTA | 1890 1891 | CD2 HIS | 129 | -0.007 -9.468 | -2.118 | 1.00 | 0.34 |
| MOTA | 1892 | HD2 HIS | 129 | 0.918 -9.447 | -2.673 | 1.00 | 0.54 |
| MOTA MOTA | 1893 | CE1 HIS HE1 HIS | 129 129 | -1.711 -8.842 | -0.936 | 1.00 | 0.31 |
| ATOM | 1894 | NE2 HIS | 129 | -2.406 -8.239 -0.597 -8.369 | -0.370 | 1.00 | 0.44 |
| MOTA | 1895 | C HIS | 129 | -0.597 -8.369 -0.614 -13.584 | -1.501 -4.277 | 1.00 | 0.28 |
| MOTA | 1896 | O HIS | 129 | -1.267 -14.568 | -4.277 -3.991 | 1.00 | 0.24 |
| ATOM | 1897 | N SER | 130 | 0.474 -13.671 | -4.999 | 1.00 | 0.28 |
| MOTA | 1898 | HN SER | 130 | 0.984 -12.862 | -5.210 | 1.00 | 0.24 |
| MOTA | 1899 | CA SER | 130 | 0.949 -14.996 | -5.498 | 1.00 | 0.29 |
| ATOM | 1900 | HA SER | 130 | 0.139 -15.710 | -5.464 | 1.00 | 0.33 |
| MOTA | 1901 | CB SER | 130 | 1.442 -14.852 | -6.938 | 1.00 | 0.32 |
| ATOM | 1902 | HB1 SER | 130 | 2.201 -14.082 | -6.982 | 1.00 | 0.31 |
| MOTA | 1903 | HB2 SER | 130 | 0.618 -14.577 | -7.576 | 1.00 | 0.35 |
| MOTA | 1904 | OG SER | 130 | 1.980 -16.092 | -7.378 | 1.00 | 0.40 |
| MOTA | 1905 | HG SER | 130 | 1.254 -16.714 | -7.469 | 1.00 | 0.97 |
| MOTA | 1906 | C SER | 130 | 2.096 -15.484 | -4.609 | 1.00 | 0.28 |
| MOTA | 1907 | O SER | 130 | 2.801 -14.696 | -4.009 | 1.00 | 0.29 |
| MOTA | 1908 | N LYS | 131 | 2.287 -16.775 | -4.514 | 1.00 | 0.30 |
| MOTA | 1909 | HN LYS | 131 | 1.705 -17.393 | -5.003 | 1.00 | 0.32 |
| MOTA | 1910 | CA LYS | 131 | 3.386 -17.310 | -3.656 | 1.00 | 0.32 |
| MOTA | 1911 | HA LYS | 131 | 3.665 -16.567 | | 1.00 | 0.34 |
| MOTA | 1912 | CB LYS | 131 | 2.903 -18.572 | | 1.00 | 0.39 |
| MOTA MOTA | 1913 1914 | HB1 LYS HB2 LYS | 131 | 3.714 -18.988 | -2.355 | 1.00 | 0.42 |
| ATOM | 1915 | HB2 LYS CG LYS | 131 131 | 2.572 -19.298 | -3.664 | 1.00 | 0.40 |
| ATOM | 1916 | HG1 LYS | 131 | . 1.743 -18.214 0.932 -17.798 | -2.003 | 1.00 | 0.45 |
| ATOM | 1917 | HG2 LYS | 131 | 2.077 -17.488 | -2.581 -1.276 | 1.00 | 0.79 |
| MOTA | 1918 | CD LYS | 131 | 1.255 -19.472 | -1.280 | 1.00 | 1.01 |
| MOTA | 1919 | HD1 LYS | 131 | 2.064 -19.890 | -0.698 | 1.00 | 1.86 |
| MOTA | 1920 | HD2 LYS | 131 | 0.921 -20.199 | -2.006 | 1.00 | 1.66 |
| MOTA | 1921 | CE LYS | 131 | 0.096 -19.108 | -0.349 | 1.00 | 1.52 |
| MOTA | 1922 | HE1 LYS | 131 | -0.788 -18.908 | -0.937 | 1.00 | 1.92 |
| MOTA | 1923 | HE2 LYS | 131 | 0.355 -18.229 | 0.222 | 1.00 | 1.93 |
| MOTA | 1924 | NZ LYS | . 131 | -0.174 -20.242 | 0.581 | 1.00 | 2.23 |
| MOTA | 1925 | HZ1 LYS | 131 | -1.103 -20.109 | 1.030 | 1.00 | 2.72 |
| MOTA | 1926 | HZ2 LYS | 131 | 0.565 -20.272 | 1.313 | 1.00 | 2.53 |
| MOTA | 1927 | HZ3 LYS | 131 | -0.174 -21.135 | 0.050 | 1.00 | 2.72 |
| MOTA | 1928 | C LYS | 131 | 4.604 -17.649 | -4.521 | 1.00 | 0.31 |
| ATOM | 1929 | O LYS | 131 | 5.612 -18.116 | -4.027 | 1.00 | 0.34 |
| MOTA | 1930 | N ASP | 132 | 4.532 -17.411 | -5.804 | 1.00 | 0.29 |
| MOTA MOTA | 1931 1932 | HN ASP | 132 | 3.717 -17.028 | -6.190 | 1.00 | 0.28 |
| ATOM | 1933 | CA ASP HA ASP | 132 · 132 | 5.703 -17.719 6.187 -18.601 | -6.674 | 1.00 | 0.30 |
| ATOM | 1934 | CB ASP | 132 | 5.225 -17.970 | -6.302 -8.108 | 1.00 | 0.32 |
| ATOM | 1935 | HB1 ASP | 132 | 4.727 -17.090 | -8.483 | 1.00 | 0.32 0.31 |
| MOTA | 1936 | HB2 ASP | 132 | 4.539 -18.804 | -8.118 | 1.00 | 0.34 |
| MOTA | 1937 | CG ASP | 132 | 6.430 -18.289 | -8.996 | 1.00 | 0.35 |
| MOTA | 1938 | OD1 ASP | 132 | 6.457 -19.371 | -9.558 | 1.00 | 1.10 |
| MOTA | 1939 | OD2 ASP | 132 | 7.306 -17.446 | -9.097 | 1.00 | 1.15 |
| MOTA | 1940 | C ASP | 132 | 6.656 -16.501 | -6.659 | 1.00 | 0.28 |
| MOTA | 1941 | O ASP | 132 | 6.226 -15.399 | -6.939 | 1.00 | 0.28 |
| MOTA | 1942 | N PRO | 133 | 7.930 -16.658 | -6.328 | 1.00 | 0.30 |
| MOTA MOTA | 1943 | CA PRO | 133 | 8.852 -15.484 | -6.296 | 1.00 | 0.31 |
| MOTA | 1944 1945 | HA PRO | 133 | 8.517 -14.766 | -5.566 | 1.00 | 0.32 |
| MOTA | 1946 | CB PRO HB1 PRO | 133 133 | 10.173 -16.097 | -5.832 | 1.00 | 0.36 |
| MOTA | 1947 | HB2 PRO | 133 | 10.441 -15.694 10.949 -15.869 | -4.867 | 1.00 | 0.36 |
| MOTA | 1948 | CG PRO | 133 | 10.007 -17.615 | -6.549 -5.721 | 1.00 | 0.41 |
| MOTA | 1949 | HG1 PRO | 133 | 10.293 -17.940 | -4.732 | 1.00 1.00 | 0.42 0.51 |
| MOTA | 1950 | HG2 PRO | 133 | 10.630 -18.103 | -6.457 | 1.00 | 0.51 |
| MOTA | 1951 | CD PRO | 133 | 8.540 -17.972 | -5.969 | 1.00 | 0.35 |
| MOTA | 1952 | HD2 PRO | 133 | 8.456 -18.679 | -6.785 | 1.00 | 0.34 |
| MOTA | 1953 | HD1 PRO | 133 | 8.091 -18.362 | -5.069 | 1.00 | 0.38 |
| MOTA | 1954 | C PRO | 133 | 9.032 -14.810 | -7.662 | 1.00 | 0.31 |
| MOTA | 1955 | O PRO | 133 | 9.496 -13.691 | -7.749 | 1.00 | 0.34 |
| MOTA | 1956 | N GLY | 134 | 8.684 -15.477 | -8.729 | 1.00 | 0.32 |
| MOTA | 1957 | HN GLY | 134 | | -8.647 | 1.00 | 0.35 |
| MOTA | 1958 | CA GLY | 134 | 8.860 -14.856 | | 1.00 | 0.34 |
| MOTA | 1959 | HA1 GLY | 134 | 9.048 -15.630 | | 1.00 | 0.37 |
| MOTA MOTA | 1960 1961 | HA2 GLY | 134 | 9.701 -14.177 | | 1.00 | 0.36 |
| ATOM | 1962 | C GLY | 134 134 | 7.598 -14.087 | | 1.00 | 0.29 |
| MOTA | 1963 | O GLY N ALA | 134 | 7.563 -13.420 6.563 -14.168 | | 1.00 | 0.29 |
| ATOM | 1964 | HN ALA | 135 | 6.607 -14.709 | -9.683 -8.867 | 1.00 | 0.27 0.28 |
| | | | | | | | |

| MOTA | 1965 | CA | ALA | 135 | 5.312 | -13.434 | -10 026 | 1.00 | 0.24 |
|--------------|--------------|----------|------------|--------------|----------------|--------------------|-------------------|--------------|--------------|
| MOTA | 1966 | HA | ALA | 135 | | -13.401 | | 1.00 | 0.25 |
| ATOM | 1967 | CB | ALA | 135 | | -14.151 | | 1.00 | 0.25 |
| MOTA | 1968 | _ | | 135 | 3.633 | -14.765 | -10.160 | 1.00 | 1.07 |
| MOTA | 1969 | | ALA | 135 | | -13.421 | | 1.00 | 1.01 |
| MOTA | 1970 | HB3 | ALA | 135 | | -14.774 | | 1.00 | 1.04 |
| ATOM | 1971 | C | ALA | 135 | | -12.007 | | 1.00 | 0.21 |
| MOTA | 1972 | 0 | ALA | 135 | | -11.760 | | 1.00 | 0.23 |
| MOTA | 1973 | N | LEU | 136 | | -11.067 | | 1.00 | 0.22 |
| MOTA MOTA | 1974 1975 | HN | LEU | 136 | | -11.286 | | 1.00 | 0.24 |
| ATOM | 1976 | CA HA | LEU | 136 136 | 4.830 | -9.660 | _ • • • • | 1.00 | 0.23 |
| ATOM | 1977 | CB | LEU | 136 | 5.842 4.279 | -9.382 | -9.427 -10.761 | 1.00 | 0.25 |
| ATOM | 1978 | HB1 | | 136 | 4.193 | | -10.761 | 1.00 | 0.25 |
| ATOM | 1979 | _ | LEU | 136 | 3.302 | | -11.064 | 1.00 | 0.27 |
| MOTA | 1980 | CG | LEU | 136 | 5.213 | | -11.980 | 1.00 | 0.26 0.26 |
| ATOM | 1981 | HG | LEU | 136 | 5.312 | | -12.368 | 1.00 | 0.29 |
| MOTA | 1982 | CD1 | LEU | 136 | 4.624 | | -13.063 | 1.00 | 0.29 |
| MOTA | 1983 | HD11 | LEU | 136 | 3.546 | | -13.030 | 1.00 | 1.06 |
| ATOM | 1984 | HD12 | LEU | 136 | 4.967 | | -14.033 | 1.00 | 1.05 |
| MOTA | 1985 | HD13 | | 136 | 4.944 | | -12.893 | 1.00 | 1.06 |
| ATOM | 1986 | CD2 | | 136 | 6.592 | -8.176 | -11.578 | 1.00 | 0.32 |
| ATOM | 1987 | HD21 | | 136 | 6.485 | -7.477 | -10.762 | 1.00 | 1.05 |
| MOTA | 1988 | HD22 | _ | 136 | 7.046 | | -12.422 | 1.00 | 1.09 |
| ATOM | | HD23 | | 136 | 7.220 | | -11.269 | 1.00 | 0.97 |
| ATOM | 1990 | C | LEU | 136 | 3.954 | -9.556 | | 1.00 | 0.25 |
| ATOM ATOM | 1991 | 0 | LEU | 136 | 4.201 | -8.761 | | 1.00 | 0.30 |
| ATOM | 1992 1993 | N | MET | 137 | 2.924 | -10.353 | | 1.00 | 0.28 |
| MOTA | 1994 | HN CA | MET | 137 | | -10.981 | | 1.00 | 0.31 |
| ATOM | 1995 | HA | MET | 137 137 | | -10.309 | | 1.00 | 0.33 |
| ATOM | 1996 | CB | MET | 137 | 1.768 | -9.283 -11.087 | -6.959 | 1.00 | 0.38 |
| ATOM | 1997 | HB1 | | 137 | | -11.136 | -7.494 | 1.00 | 0.42 |
| MOTA | 1998 | HB2 | MET | 137 | | -12.089 | -6.615 -7.803 | 1.00 | 0.57 |
| ATOM | 1999 | CG | MET | 137 | | -10.391 | | 1.00 1.00 | 0.50 0.58 |
| MOTA | 2000 | HG1 | | 137 | | -10.975 | -8.875 | 1.00 | 1.13 |
| MOTA | 2001 | HG2 | - | 137 | | -10.311 | -9.494 | 1.00 | 1.22 |
| ATOM | 2002 | SD | MET | 137 | -0.551 | | | 1.00 | 0.83 |
| MOTA | 2003 | CE | MET | 137 · | -2.048 | | -7.194 | 1.00 | 0.39 |
| MOTA | 2004 | HE1 | | 137 | -2.231 | - | -6.426 | 1.00 | 1.14 |
| ATOM | 2005 | | MET | 137 | -1.927 | -10.151 | | 1.00 | 1.07 |
| MOTA | 2006 | HE3 | MET | 137 | -2.885 | -9.212 | -7.872 | 1.00 | 1.06 |
| ATOM | 2007 | C | MET | 137 | | -10.925 | -5.951 | 1.00 | 0.27 |
| ATOM | 2008 | 0 | MET | 137 | | -11.287 | -4.990 | 1.00 | 0.28 |
| MOTA | 2009 | N | PHE | 138 | | -11.042 | -5.964 | 1.00 | 0.25 |
| MOTA MOTA | 2010 2011 | HN | PHE | 138 | | -10.741 | -6.743 | 1.00 | 0.28 |
| ATOM | 2012 | CA HA | PHE | 138 | | -11.628 | -4.785 | 1.00 | 0.23 |
| ATOM | 2012 | CB | PHE | 138 138 | _ | -12.557 | -4.534 | 1.00 | 0.26 |
| MOTA | 2014 | HB1 | PHE | 138 | | -11.877 -10.945 | -5.152 | 1.00 | 0.25 |
| ATOM | 2015 | | PHE | 138 | | -10.945 | -5.104 | 1.00 | 0.24 |
| MOTA | 2016 | CG | PHE | 138 | | -12.873 | -6.156 -4.194 | 1.00 | 0.27 |
| ATOM | 2017 | | PHE | 138 | | -14.184 | -4.113 | 1.00 | 0.28 0.32 |
| ATOM | 2018 | | PHE | 138 | | -14.490 | -4.731 | 1.00 | 0.32 |
| MOTA | 2019 | CD2 | PHE | 138 | | -12.486 | -3.392 | 1.00 | 0.30 |
| MOTA | 2020 | HD2 | PHE | 138 | | -11.481 | -3.455 | 1.00 | 0.30 |
| MOTA | 2021 | | PHE | 138 | _ | -15.100 | -3.230 | 1.00 | 0.38 |
| MOTA | 2022 | | PHE | 138 | 6.500 | -16.109 | -3.168 | 1.00 | 0.42 |
| MOTA | 2023 | | PHE | | | -13.404 | -2.511 | 1.00 | 0.36 |
| ATOM | 2024 | HE2 | PHE | 138 | | -13.104 | -1.894 | 1.00 | 0.39 |
| MOTA | 2025 | CZ | PHE | 138 | | -14.710 | -2.430 | 1.00 | 0.39 |
| MOTA | 2026 | HZ | PHE | 138 | | -15.417 | -1.749 | 1.00 | 0.44 |
| MOTA MOTA | 2027 | C | PHE | 138 | | -10.615 | -3.615 | 1.00 | 0.20 |
| ATOM | 2028 | 0 | PHE | 138 | 4.874 | -9.447 | -3.808 | 1.00 | 0.22 |
| ATOM | 2029 2030 | N C2 | PRO | 139 | | -11.019 | -2.421 | 1.00 | 0.22 |
| ATOM | 2031 | CA HA | PRO | 139 | | -10.048 | ~1.291 | 1.00 | 0.25 |
| ATOM | 2032 | CB | PRO PRO | 139 139 | 3.262 | -9.340 | -1.509 | 1.00 | 0.27 |
| ATOM | 2032 | HB1 | PRO | 139 | | -10.936 -10.638 | -0.127 | 1.00 | 0.31 |
| ATOM | 2034 | HB2 | PRO | 139 | | -10.638 | 0.199 | 1.00 | 0.38 |
| ATOM | 2035 | CG | PRO | 139 | | -12.392 | 0.691 -0.597 | 1.00 | 0.42 |
| MOTA | 2036 | HG1 | | 139 | | -12.812 | -0.397 | 1.00 | 0.33 0.41 |
| ATOM | 2037 | HG2 | PRO | 139 | | -12.961 | -0.336 | 1.00 | 0.41 |
| MOTA | 2038 | CD | PRO | 139 | | -12.435 | -2.102 | 1.00 | 0.27 |
| MOTA | 2039 | HD2 | PRO | 139 | | -13.100 | -2.318 | 1.00 | 0.28 |
| MOTA | 2040 | HD1 | PRO | 139 | 2.946 | -12.732 | -2.637 | 1.00 | 0.30 |
| MOTA | 2041 | C | טמם | 120 | E 777 | -0 305 | 0 026 | 1 00 | 0.50 |

| MOTA | 2042 | 0 1 | PRO | 139 | | 5.302 | -8.351 | -0.173 | 1.00 | 0.44 |
|------|--------|-------------|-----|-----|---|--------|-------------|---------|------|------|
| ATOM | 2043 | | | | | | | _ | | |
| | | | ILE | 140 | | 6.467 | -9.726 | -1.437 | 1.00 | 0.24 |
| MOTA | 2044 | | ILE | 140 | | 6.474 | -10.500 | -2.038 | 1.00 | 0.37 |
| MOTA | 2045 | CA 3 | ILE | 140 | | 7.749 | -9.031 | -1.094 | 1.00 | 0.23 |
| MOTA | 2046 | HA I | ILE | 140 | | 7.572 | -8.308 | -0.312 | 1.00 | 0.24 |
| MOTA | 2047 | | ILE | 140 | | | -10.054 | | | |
| | | | | | | | | -0.600 | 1.00 | 0.25 |
| MOTA | 2048 | | ILE | 140 | | | -10.770 | -1.379 | 1.00 | 0.25 |
| MOTA | 2049 | CG1 | ILE | 140 | | 8.207 | -10.768 | 0.632 | 1.00 | 0.29 |
| MOTA | 2050 | HG11 | ILE | 140 | | 7.246 | -11.196 | 0.384 | 1.00 | 0.32 |
| MOTA | | HG12 | | 140 | | | -10.055 | 1.434 | | |
| | | | | | | | | | 1.00 | 0.33 |
| MOTA | 2052 | | ILE | 140 | | 10.070 | | -0.214 | 1.00 | 0.26 |
| MOTA | 2053 | HG21 | ILE | 140 | | 9.850 | -8.567 | 0.517 | 1.00 | 1.04 |
| MOTA | 2054 | HG22 | ILE | 140 | | 10.505 | -8.876 | -1.090 | 1.00 | 1.06 |
| MOTA | 2055 | | ILE | 140 | | | -10.040 | 0.207 | 1.00 | 1.04 |
| ATOM | 2056 | | ILE | 140 | | | -11.883 | | | |
| | - | | | | | | | 1.082 | 1.00 | 0.30 |
| ATOM | | HD11 : | | 140 | | | -12.250 | 0.236 | 1.00 | 1.08 |
| MOTA | 2058 | HD12 | ILE | 140 | | 8.582 | -12.691 | 1.511 | 1.00 | 0.98 |
| MOTA | 2059 | HD13 : | ILE | 140 | | 9.838 | -11.495 | 1.824 | 1.00 | 1.08 |
| MOTA | 2060 | | ILE | 140 | | 8.284 | | -2.329 | 1.00 | |
| | | | | | | | | | | 0.22 |
| MOTA | 2061 | | ILE | 140 | | 8.265 | -8.817 | -3.429 | 1.00 | 0.22 |
| MOTA | 2062 | N | TYR | 141 | | 8.745 | -7.092 | -2.150 | 1.00 | 0.21 |
| ATOM | 2063 | HN ' | TYR | 141 | | 8.736 | -6.696 | -1.254 | 1.00 | 0.22 |
| MOTA | 2064 | | TYR | 141 | | 9.265 | -6.303 | -3.304 | 1.00 | 0.21 |
| ATOM | 2065 | | | | | | | | | |
| | | | TYR | 141 | | 8.560 | -6.348 | -4.120 | 1.00 | 0.20 |
| MOTA | 2066 | | TYR | 141 | | 9.444 | -4.847 | -2.865 | 1.00 | 0.21 |
| MOTA | 2067 | HB1 ' | TYR | 141 | | 10.050 | -4.810 | -1.972 | 1.00 | 0.22 |
| MOTA | 2068 | HB2 | TYR | 141 | | 8.476 | -4.413 | -2.661 | 1.00 | 0.22 |
| MOTA | 2069 | | TYR | 141 | | | | | _ | |
| - | | | | | | 10.122 | -4.066 | -3.962 | 1.00 | 0.23 |
| MOTA | 2070 | CD1 | | 141 | | 11.515 | -4.104 | -4.089 | 1.00 | 0.25 |
| MOTA | 2071 | HD1 | TYR | 141 | | 12.104 | -4.697 | -3.404 | 1.00 | 0.26 |
| MOTA | 2072 | CD2 | TYR | 141 | | 9.359 | -3.298 | -4.848 | 1.00 | 0.24 |
| MOTA | 2073 | | TYR | 141 | | 8.284 | -3.268 | -4.750 | 1.00 | |
| _ | 2074 | | | | | _ | | | | 0.25 |
| MOTA | | | TYR | 141 | | 12.146 | -3.376 | -5.103 | 1.00 | 0.28 |
| MOTA | 2075 | | TYR | 141 | | 13.221 | -3.405 | -5.201 | 1.00 | 0.32 |
| ATOM | 2076 | CE2 | TYR | 141 | | 9.989 | -2.569 | -5.862 | 1.00 | 0.27 |
| ATOM | 2077 | HE2 | TYR | 141 | | 9.401 | -1.975 | -6.544 | 1.00 | 0.30 |
| MOTA | 2078 | | TYR | 141 | | | | | | |
| | | _ | | | | 11.383 | -2.608 | -5.990 | 1.00 | 0.29 |
| MOTA | 2079 | | TYR | 141 | | 12.005 | | -6.991 | 1.00 | 0.33 |
| ATOM | 2080 | HH | TYR | 141 | | 12.781 | -2.385 | -7.269 | 1.00 | 0.90 |
| ATOM | 2081 | С | TYR | 141 | | 10.615 | -6.864 | -3.761 | 1.00 | 0.22 |
| MOTA | 2082 | | TYR | 141 | | 11.522 | | -2.973 | _ | |
| ATOM | 2083 | | | | | | | | 1.00 | 0.23 |
| | | | THR | 142 | | 10.750 | | -5.035 | 1.00 | 0.22 |
| MOTA | 2084 | HN | THR | 142 | | 10.002 | -6.968 | -5.648 | 1.00 | 0.22 |
| MOTA | 2085 | ÇA | THR | 142 | | 12.035 | -7.675 | -5.563 | 1.00 | 0.24 |
| ATOM | 2086 | HA | THR | 142 | | 12.835 | | -4.874 | 1.00 | 0.25 |
| ATOM | 2087 | | THR | 142 | | 11.917 | | | | |
| | | | | | | | | -5.723 | 1.00 | 0.25 |
| ATOM | 2088 | | THR | 142 | | 11.645 | _ | -4.777 | 1.00 | 0.26 |
| MOTA | 2089 | OG1 | THR | 142 | | 13.165 | -9.720 | -6.152 | 1.00 | 0.29 |
| MOTA | 2090 | HG1 | THR | 142 | | 13.274 | -9.505 | -7.081 | 1.00 | 0.97 |
| MOTA | 2091 | CG2 | THR | 142 | | 10.840 | | -6.760 | 1.00 | 0.25 |
| MOTA | | | THR | 142 | | | | | | |
| | | | | | | 10.577 | | -6.691 | 1.00 | 1.04 |
| MOTA | 2093 | | THR | 142 | | 11.217 | | -7.749 | 1.00 | 1.05 |
| MOTA | 2094 | HG23 | THR | 142 | | 9.965 | -8.913 | -6.570 | 1.00 | 1.06 |
| ATOM | · 2095 | C | THR | 142 | | 12.339 | -7.040 | -6.924 | 1.00 | 0.23 |
| MOTA | 2096 | 0 | THR | 142 | | 11.454 | | -7.724 | 1.00 | 0.23 |
| ATOM | 2097 | | TYR | 143 | - | 13.586 | | | | |
| ATOM | 2098 | | | | - | | | -7.195 | 1.00 | 0.25 |
| | | | TYR | 143 | | 14.285 | | -6.538 | 1.00 | 0.27 |
| MOTA | 2099 | | TYR | 143 | | 13.948 | -6.144 | -8.506 | 1.00 | 0.26 |
| MOTA | 2100 | HA | TYR | 143 | | 13.174 | -5.452 | -8.804 | 1.00 | 0.25 |
| MOTA | 2101 | CB | TYR | 143 | | 15.277 | | -8.370 | 1.00 | 0.29 |
| ATOM | 2102 | | TYR | 143 | | 16.072 | | | | |
| ATOM | 2103 | _ | | | | | | -8.190 | 1.00 | 0.33 |
| | | | TYR | 143 | | 15.217 | | -7.542 | 1.00 | 0.30 |
| ATOM | 2104 | | TYR | 143 | | 15.563 | -4.633 | -9.642 | 1.00 | 0.27 |
| MOTA | 2105 | CD1 | TYR | 143 | | 14.931 | -3.406 | -9.880 | 1.00 | 0.25 |
| MOTA | 2106 | HD1 | | 143 | | 14.234 | | | 1.00 | 0.26 |
| ATOM | 2107 | CD2 | | 143 | | | _ | | | |
| | | | | | | 16.466 | | -10.581 | 1.00 | 0.31 |
| MOTA | 2108 | HD2 | | 143 | | 16.954 | · · · - • - | -10.398 | 1.00 | 0.35 |
| MOTA | 2109 | CE1 | | 143 | | 15.201 | -2.695 | -11.055 | 1.00 | 0.26 |
| MOTA | 2110 | HE1 | TYR | 143 | | 14.713 | | -11.238 | 1.00 | 0.28 |
| ATOM | 2111 | CE2 | | 143 | | 16.735 | | -11.756 | 1.00 | |
| ATOM | 2112 | HE2 | | | | | | | | 0.31 |
| | | | | 143 | | 17.432 | | -12.480 | 1.00 | 0.36 |
| MOTA | 2113 | | TYR | 143 | | 16.103 | _ | -11.994 | 1.00 | 0.28 |
| MOTA | 2114 | OH | TYR | 143 | | 16.369 | -2.509 | -13.152 | 1.00 | 0.30 |
| MOTA | 2115 | HH | TYR | 143 | | 17.068 | | -13.624 | 1.00 | 0.95 |
| MOTA | 2116 | C | TYR | 143 | | 14.080 | | -9.563 | 1.00 | 0.27 |
| MOTA | 2117 | Ö | TYR | 143 | | 14.552 | | | | |
| ATOM | 2118 | N | THR | 144 | | 13 660 | | -9.283 | 1.00 | 0.31 |
| | | | | | | | | | | |

| > 000M | 2110 | | MIID | • • • | 42 222 | | 40 000 | - 00 | |
|---------------------|-------|------|------|-------|--------|---------|---------|-------|------|
| MOTA | 2119 | HN | THR | 144 | 13.277 | -6.096 | | 1.00 | 0.32 |
| MOTA | 2120 | CA | THR | 144 | 13.753 | -8.008 | -11.847 | 1.00 | 0.32 |
| MOTA | 2121 | HA | THR | 144 | 14.479 | -8.758 | -11.573 | 1.00 | 0.35 |
| MOTA | 2122 | CB | THR | 144 | 12.385 | -8.666 | | 1.00 | 0.37 |
| MOTA | 2123 | | THR | 144 | | | | | |
| | | HB | | | 11.922 | -8.814 | | 1.00 | 0.84 |
| MOTA | 2124 | OG1 | THR | 144 | 12.549 | -9.918 | -12.683 | 1.00 | 1.00 |
| MOTA | 2125 | HG1 | THR | 144 | 13.280 | -9.836 | -13.301 | 1.00 | 1.42 |
| MOTA | 2126 | CG2 | THR | 144 | 11.499 | -7.757 | -12.882 | 1.00 | 0.82 |
| MOTA | | HG21 | THR | 144 | 10.461 | -7.991 | | 1.00 | 1.51 |
| ATOM | | HG22 | THR | 144 | 11.724 | | | | |
| | | | | | | -7.911 | | 1.00 | 1.24 |
| MOTA | | HG23 | THR | 144 | 11.687 | | -12.622 | 1.00 | 1.49 |
| MOTA | 2130 | C | THR | 144 | 14.169 | -7.351 | -13.165 | 1.00 | 0.34 |
| MOTA | 2131 | 0 | THR | 144 | 13.922 | -6.183 | -13.392 | 1.00 | 0.32 |
| MOTA | 2132 | N | GLY | 145 | 14.789 | | -14.043 | 1.00 | 0.43 |
| ATOM | 2133 | HN | GLY | 145 | 14.971 | | | | _ |
| MOTA | 2134 | | | | | | -13.846 | 1.00 | 0.49 |
| | | CA | GLY | 145 | 15.205 | | -15.350 | 1.00 | 0.49 |
| MOTA | 2135 | HA1 | GLY | 145 | 15.842 | -8.207 | -15.872 | 1.00 | 0.57 |
| MOTA | 2136 | HA2 | GLY | 145 | 15.742 | -6.587 | -15.178 | 1.00 | 0.50 |
| MOTA | 2137 | C | GLY | 145 | 13.957 | -7.233 | -16.191 | 1.00 | 0.47 |
| ATOM | 2138 | 0 | GLY | 145 | 13.331 | -8.138 | -16.706 | 1.00 | 0.53 |
| ATOM | 2139 | N | LYS | 146 | 13.583 | | -16.322 | 1.00 | 0.46 |
| MOTA | 2140 | HN | LYS | 146 | 14.097 | | | | |
| ATOM | 2141 | | | | | | -15.889 | 1.00 | 0.48 |
| | | CA | LYS | 146 | 12.367 | | -17.116 | 1.00- | 0.49 |
| MOTA | 2142 | HA | LYS | 146 | 11.578 | -6.350 | -16.876 | 1.00 | 0.51 |
| MOTA | 2143 | CB | LYS | 146 | 11.911 | -4.235 | -16.764 | 1.00 | 0.52 |
| MOTA | 2144 | HB1 | LYS | 146 | 10.973 | -4.032 | -17.254 | 1.00 | 0.58 |
| MOTA | 2145 | HB2 | LYS | 146 | 12.657 | | -17.103 | 1.00 | 0.57 |
| MOTA | 2146 | CG | LYS | 146 | 11.744 | | -15.238 | | |
| MOTA | 2147 | | LYS | 146 | | | | 1.00 | 0.55 |
| | | | | | 12.690 | | -14.798 | 1.00 | 0.83 |
| MOTA | 2148 | | LYS | 146 | 11.442 | | -14.849 | 1.00 | 1.14 |
| MOTA | 2149 | CD | LYS | 146 | 10.684 | -3.077 | -14.854 | 1.00 | 1.23 |
| ATOM | 2150 | HD1 | LYS | 146 | 10.308 | -3.309 | -13.871 | 1.00 | 1.78 |
| MOTA | 2151 | HD2 | LYS | 146 | 9.865 | | -15.556 | 1.00 | 1.79 |
| MOTA | 2152 | CE | LYS | 146 | 11.298 | | -14.828 | | - |
| MOTA | 2153 | | LYS | 146 | | | | 1.00 | 2.01 |
| MOTA | 2154 | | | | 11.615 | | -13.822 | 1.00 | 2.47 |
| | | | LYS | 146 | 10.556 | | -15.143 | 1.00 | 2.39 |
| ATOM | 2155 | NZ | LYS | 146 | 12.468 | -1.601 | -15.745 | 1.00 | 2.91 |
| MOTA | 2156 | HZ1 | LYS | 146 | 12.847 | -0.633 | -15.750 | 1.00 | 3.39 |
| MOTA | 2157 | HZ2 | LYS | 146 | 12.170 | | -16.707 | 1.00 | 3.28 |
| MOTA | 2158 | | LYS | 146 | 13.205 | | -15.420 | 1.00 | 3.27 |
| ATOM | 2159 | C | LYS | 146 | | | | | |
| ATOM | 2160 | | | | 12.677 | | -18.613 | 1.00 | 0.59 |
| | _ | 0 | LYS | 146 | 11.845 | | -19.444 | 1.00 | 1.16 |
| ATOM | 2161 | N | SER | 147 | 13.868 | -6.131 | -18.967 | 1.00 | 0.78 |
| MOTA | 2162 | HN | SER | 147 | 14.530 | -6.366 | -18.283 | 1.00 | 1.26 |
| MOTA | 2163 | CA | SER | 147 | 14.226 | | -20.413 | 1.00 | 0.87 |
| MOTA | 2164 | HA | SER | 147 | 14.141 | | -20.859 | 1.00 | 1.03 |
| ATOM | 2165 | CB | SER | 147 | 15.667 | | | | |
| MOTA | 2166 | HB1 | | | | | -20.554 | 1.00 | 0.95 |
| | | | SER | 147 | 15.798 | | -21.530 | 1.00 | 1.42 |
| MOTA | 2167 | HB2 | | 147 | 15.871 | -7.445 | -19.794 | 1.00 | 1.34 |
| MOTA | 2158 | OG | SER | 147 | 16.561 | -5.616 | -20.395 | 1.00 | 1.71 |
| MOTA | 2169 | HG | SER | 147 | 17.097 | -5.555 | -21.190 | 1.00 | 2.16 |
| MOTA | 2170 | C | SER | 147 | 13.288 | | -21.138 | 1.00 | 0.79 |
| MOTA | 2171 | 0 | SER | 147 | 12.747 | | -22.178 | | |
| ATOM | 2172 | N | HIS | 148 | | | | 1.00 | 1.40 |
| MOTA | 2173 | | | | 13.098 | | -20.605 | 1.00 | 0.66 |
| MOTA | | HN | HIS | 148 | 13.551 | | -19.768 | 1.00 | 1.10 |
| | 2174 | CA | HIS | 148 | 12.199 | | -21.272 | 1.00 | 0.65 |
| MOTA | 2175 | HA | HIS | 148 | 11.629 | | -22.048 | 1.00 | 0.74 |
| MOTA | 2176 | CB | HIS | 148 | 13.041 | -10.479 | -21.887 | 1.00 | 0.79 |
| MOTA | 2177 | HB1 | HIS | 148 | | -11.312 | | 1.00 | 1.14 |
| MOTA | 2178 | | HIS | 148 | | -10.801 | | 1.00 | 1.30 |
| MOTA | 2179 | CG | HIS | 148 | | -9.980 | | | |
| MOTA | 2180 | | HIS | | | | | 1.00 | 1.66 |
| | | | | 148 | | -9.116 | | 1.00 | 2.52 |
| MOTA | 2181 | | HIS | 148 | 12.200 | | -23.934 | 1.00 | 2.81 |
| MOTA | 2182 | | HIS | 148 | | -10.226 | | 1.00 | 2.62 |
| MOTA | 2183 | | HIS | 148 | 15.715 | -10.867 | -23.206 | 1.00 | 3.00 |
| MOTA | 2184 | CEl | HIS | 148 | 13.970 | | -25.020 | 1.00 | 3.46 |
| MOTA | 2185 | HE1 | HIS | 148 | 13.759 | | -25.863 | 1.00 | 4.33 |
| MOTA | 2186 | NE2 | | 148 | 15.123 | | -24.846 | | |
| MOTA | 2187 | C | HIS | 148 | | | | 1.00 | 3.55 |
| MOTA | 2188 | | | | 11.238 | | -20.249 | 1.00 | 0.55 |
| | | 0 | HIS | 148 | | -11.064 | | 1.00 | 0.60 |
| MOTA | 2189 | N | PHE | 149 | 10.978 | | -19.167 | 1.00 | 0.57 |
| MOTA | 2190 | HN | PHE | 149 | 11.392 | -8.417 | -19.021 | 1.00 | 0.73 |
| MOTA | 2191 | CA | PHE | 149 | 10.060 | | -18.145 | 1.00 | 0.48 |
| MOTA | 2192 | HA | PHE | 149 | | -10.849 | | 1.00 | 0.51 |
| MOTA | 2193 | CB | PHE | 149 | 10.022 | | -16.911 | | |
| MOTA | 2194 | HB1 | PHE | 149 | 9.603 | | | 1.00 | 0.44 |
| MOTA | 2195 | | PHE | | | | -17.177 | 1.00 | 0.44 |
| saa Vi i | E 173 | nb2 | FRE | 149 | 11.023 | -x.831 | -16 530 | 1 11 | U 46 |
| | | | | | | | | | |

| MOTA | 2196 | CG | PHE | 149 | 9.161 -9.615 -15.851 1.00 0.40 |
|--------------|--------------|--------------|-----|------------|--|
| MOTA | 2197 | | PHE | 149 | 7.766 -9.507 -15.919 1.00 0.36 |
| MOTA | 2198 | _ | PHE | 149 | 7.305 -8.956 -16.726 1.00 0.38 |
| MOTA | 2199 | | PHE | 149 | 9.757 -10.328 -14.804 1.00 0.42 |
| MOTA | 2200 | HD2 | PHE | 149 | 10.832 -10.412 -14.750 1.00 0.48 |
| MOTA | 2201 | CE1 | PHE | 149 | 6.969 -10.112 -14.941 1.00 0.35 |
| MOTA | 2202 | | PHE | 149 | 5.894 -10.031 -14.996 1.00 0.37 |
| MOTA | 2203 | | PHE | 149 | 8.958 -10.932 -13.825 1.00 0.40 |
| MOTA | 2204 | | PHE | 149 | 9.417 -11.482 -13.016 1.00 0.45 |
| MOTA | 2205 | CZ | PHE | 149 | 7.564 -10.825 -13.894 1.00 0.37 |
| MOTA | 2206 | HZ | PHE | 149 | 6.948 -11.291 -13.140 1.00 0.38 |
| MOTA | 2207 | C | PHE | 149 | 8.641 -9.993 -18.706 1.00 0.43 |
| MOTA MOTA | 2208 2209 | 0 N | PHE | 149 | 8.080 -9.044 -19.217 1.00 0.45 |
| MOTA | 2210 | N HN | MET | 150 150 | 8.050 -11.153 -18.575 1.00 0.43 |
| MOTA | 2211 | CA | MET | 150 | 8.523 -11.888 -18.133 1.00 0.50 6.651 -11.357 -19.051 1.00 0.39 |
| ATOM | 2212 | HA | MET | 150 | 6.651 -11.357 -19.051 1.00 0.39 6.189 -10.400 -19.245 1.00 0.38 |
| ATOM | 2213 | CB | MET | 150 | 6.632 -12.207 -20.328 1.00 0.44 |
| MOTA | 2214 | HB1 | MET | 150 | 5.610 -12.374 -20.632 1.00 0.45 |
| MOTA | 2215 | HB2 | MET | 150 | 7.109 -13.157 -20.134 1.00 0.47 |
| MOTA | 2216 | CG | MET | 150 | 7.381 -11.477 -21.446 1.00 0.50 |
| MOTA | 2217 | HG1 | MET | 150 | 8.401 -11.831 -21.485 1.00 0.98 |
| MOTA | 2218 | | MET | 150 | 7.376 -10.415 -21.253 1.00 0.86 |
| ATOM | 2219 | SD | MET | 150 | 6.571 -11.806 -23.033 1.00 1.32 |
| MOTA | 2220 | CE | MET | 150 | 7.378 -13.384 -23.393 1.00 2.23 |
| MOTA | 2221 | HE1 | MET | 150 | 7.326 -14.022 -22.521 1.00 2.66 |
| MOTA | 2222 | HE2 | MET | 150 | 8.411 -13.211 -23.647 1.00 2.74 |
| MOTA MOTA | 2223 2224 | | MET | 150 | 6.879 -13.861 -24.225 1.00 2.74 |
| MOTA | 2225 | C | MET | 150 | 5.877 -12.071 -17.943 1.00 0.32 |
| MOTA | 2226 | N | LEU | 150 151 | 6.435 -12.837 -17.183 1.00 0.32 4.605 -11.819 -17.827 1.00 0.28 |
| ATOM | 2227 | HN | LEU | 151 | 4.605 -11.819 -17.827 1.00 0.28 4.169 -11.188 -18.437 1.00 0.30 |
| ATOM | 2228 | CA | LEU | 151 | 3.821 -12.478 -16.746 1.00 0.24 |
| ATOM | 2229 | HA | LEU | 151 | 4.120 -12.064 -15.803 1.00 0.24 |
| MOTA | 2230 | CB | LEU | 151 | 2.327 -12.212 -16.966 1.00 0.24 |
| ATOM | 2231 | | LEU | 151 | 1.765 -12.626 -16.145 1.00 0.25 |
| MOTA | 2232 | | | 151 | 2.012 -12.680 -17.887 1.00 0.28 |
| MOTA | 2233 | CG | LEU | 151 | 2.061 -10.703 -17.047 1.00 0.28 |
| MOTA | 2234 | HG | LEU | 151 | 2.900 -10.208 -17.512 1.00 0.52 |
| MOTA | 2235 | | LEU | 151 | 0.804 -10.457 -17.881 1.00 0.35 |
| MOTA | | HD11 | | | 0.506 -9.424 -17.788 1.00 1.07 |
| ATOM | 2237 | HD12 | | 151 | 0.007 -11.095 -17.526 1.00 1.02 |
| MOTA | 2238 | | | 151 | 1.009 -10.682 -18.917 1.00 1.17 |
| ATOM | 2239 | | LEU | 151 | 1.848 -10.140 -15.638 1.00 0.46 |
| MOTA MOTA | | HD21 HD22 | | 151 | 2.078 -9.084 -15.635 1.00 1.14 |
| MOTA | | | | 151 151 | 2.495 -10.650 -14.941 1.00 1.16 |
| ATOM | 2243 | C | LEU | 151 | 0.820 -10.284 -15.345 1.00 1.11 4.076 -14.004 -16.794 1.00 0.24 |
| ATOM | 2244 | Ö | LEU | 151 | 4.076 -14.004 -16.794 1.00 0.24 3.879 -14.613 -17.826 1.00 0.28 |
| MOTA | 2245 | N | PRO | 152 | 4.504 -14.641 -15.711 1.00 0.22 |
| MOTA | 2246 | CA | PRO | 152 | 4.748 -16.112 -15.751 1.00 0.23 |
| MOTA | 2247 | HA | PRO | 152 | 5.480 -16.354 -16.503 1.00 0.24 |
| MOTA | 2248 | CB | PRO | 152 | 5.323 -16.404 -14.364 1.00 0.24 |
| MOTA | 2249 | HB1 | PRO | 152 | 6.361 -16.686 -14.453 1.00 0.29 |
| MOTA | 2250 | HB2 | | 152 | 4.766 -17.208 -13.903 1.00 0.26 |
| MOTA | 2251 | CG | PRO | 152 | 5.209 -15.141 -13.507 1.00 0.32 |
| MOTA | 2252 | | PRO | 152 | 6.166 -14.917 -13.061 1.00 0.44 |
| MOTA | 2253 | | PRO | 152 | 4.473 -15.295 -12.730 1.00 0.41 |
| MOTA MOTA | 2254 2255 | | PRO | 152 | 4.778 -13.976 -14.402 1.00 0.25 |
| MOTA | 2256 | HD2 HD1 | _ | 152 | 3.886 -13.507 -14.008 1.00 0.25 |
| MOTA | 2257 | C | PRO | 152 152 | 5.581 -13.263 -14.503 1.00 0.27 |
| MOTA | 2258 | Õ | PRO | 152 | 3.462 -16.915 -15.974 1.00 0.21 2.378 -16.371 -16.038 1.00 0.20 |
| ATOM | 2259 | N | ASP | 153 | |
| ATOM | 2260 | | ASP | 153 | 3.582 -18.209 -16.090 1.00 0.23 4.468 -18.622 -16.031 1.00 0.25 |
| MOTA | 2261 | CA | ASP | 153 | 2.380 -19.063 -16.304 1.00 0.23 |
| MOTA | 2262 | | ASP | 153 | 1.890 -18.772 -17.221 1.00 0.23 |
| MOTA | 2263 | CB | ASP | 153 | 2.813 -20.526 -16.401 1.00 0.25 |
| MOTA | 2254 | HB1 | ASP | | 1.943 -21.163 -16.363 1.00 0.26 |
| MOTA | 2265 | HB2 | ASP | 153 | 3.470 -20.762 -15.576 1.00 0.26 |
| MOTA | 2266 | | | 153 | 3.550 -20.752 -17.722 1.00 0.27 |
| MOTA | 2267 | | ASP | 153 | 4.768 -20.687 -17.717 1.00 1.08 |
| MOTA | 2268 | | ASP | | 2.884 -20.994 -18.715 1.00 1.14 |
| MOTA | 2269 | | ASP | 153 | 1.409 -18.899 -15.133 1.00 0.21 |
| MOTA MOTA | 2270 2271 | - | ASP | 153 | 0.208 -18.858 -15.310 1.00 0.21 |
| MOTA | 2271 | | ASP | 154 154 | 1.919 -18.820 -13.935 1.00 0.21 |
| | a. a. s 1. | - IA | | . 74 | 7 991 -10 044 -13 013 1 00 0 00 |

| MOTA | 2273 | CA | ASP | 154 | 1.025 -18.678 -12.752 1.00 0.3 | 21 |
|------|------|-----------|------|------------|---|-----------|
| MOTA | 2274 | HA | ASP | 154 | 0.431 -19.572 -12.641 1.00 0.3 | 22 |
| MOTA | 2275 | CB | ASP | 154 | 1.880 -18.474 -11.496 1.00 0.3 | 23 |
| MOTA | 2276 | HB1 | ASP | 154 | 2.466 -17.572 -11.602 1.00 0.3 | |
| ATOM | 2277 | HB2 | ASP | 154 | 2.541 -19.319 -11.370 1.00 0.3 | |
| ATOM | 2278 | CG | ASP | 154 | 0.975 -18.347 -10.267 1.00 0.3 | |
| ATOM | 2279 | | ASP | 154 | 1.276 -18.982 -9.269 1.00 1.3 | |
| ATOM | 2280 | | ASP | 154 | | |
| | | | | | 0.004 -17.613 -10.340 1.00 1.0 | |
| ATOM | 2281 | C | ASP | 154 | 0.102 -17.473 -12.943 1.00 0.3 | - |
| ATOM | 2282 | 0 | ASP | 154 | -1.095 -17.564 -12.759 1.00 0.3 | 19 |
| ATOM | 2283 | N | ASP | 155 | 0.645 -16.345 -13.303 1.00 0.3 | 19 |
| MOTA | 2284 | HN | ASP | 155 | 1.613 -16.288 -13.443 1.00 0. | |
| MOTA | 2285 | CA | ASP | 155 | -0.210 -15.140 -13.496 1.00 0. | |
| ATOM | 2286 | HA | ASP | 155 | | |
| ATOM | 2287 | CB | ASP | 155 | | |
| | | | | | 0.683 -13.909 -13.653 1.00 0.3 | |
| ATOM | 2288 | | ASP | 155 | 0.087 -13.067 -13.969 1.00 0.3 | |
| ATOM | 2289 | | | 155 | 1.443 -14.113 -14.393 1.00 0.3 | 22 |
| MOTA | 2290 | CG | ASP | 155 | 1.351 -13.588 -12.315 1.00 0.5 | 24 |
| MOTA | 2291 | OD1 | ASP | 155 | 2.355 -12.896 -12.327 1.00 1. | 07 |
| MOTA | 2292 | OD2 | ASP | 155 | 0.845 -14.038 -11.300 1.00 1. | |
| ATOM | 2293 | С | ASP | 155 | -1.087 -15.300 -14.744 1.00 0. | |
| ATOM | 2294 | Ö | ASP | 155 | | |
| ATOM | 2295 | N | VAL | 156 | | |
| | | | | | -0.555 -15.850 -15.802 1.00 0. | - |
| MOTA | 2296 | HN | VAL | 156 | 0.379 -16.147 -15.787 1.00 0. | |
| MOTA | 2297 | CA | VAL | 156 | -1.372 -16.013 -17.041 1.00 0. | 21 |
| MOTA | 2298 | HA | VAL | 156 | -1.726 -15.044 -17.362 1.00 0.3 | 22 |
| ATOM | 2299 | CB | VAL | 156 | -0.519 -16.630 -18.148 1.00 0. | 23 |
| ATOM | 2300 | HB | VAL | 156 | -0.034 -17.521 -17.776 1.00 0. | |
| MOTA | 2301 | | VAL | 156 | -1.416 -16.995 -19.333 1.00 0. | |
| ATOM | | HG11 | | 156 | | |
| ATOM | | HG12 | | | | |
| | | | | 156 | -1.747 -18.018 -19.235 1.00 1. | 05 |
| ATOM | 2304 | - · · · - | | 156 | -0.861 -16.882 -20.253 1.00 1. | 05 |
| MOTA | 2305 | | VAL | 156 | 0.535 -15.618 -18.600 1.00 0. | 26 |
| MOTA | 2306 | HG21 | VAL | 156 | 0.990 -15.162 -17.733 1.00 1. | 07 |
| ATOM | 2307 | HG22 | VAL | 156 | 0.067 -14.856 -19.204 1.00 1. | |
| ATOM | 2308 | HG23 | | 156 | 1.293 -16.123 -19.180 1.00 1. | |
| MOTA | 2309 | | VAL | 156 | | |
| ATOM | 2310 | | | | | _ |
| | | _ | VAL | 156 | -3.694 -16.615 -17.107 1.00 0. | |
| MOTA | 2311 | | GLN | 157 | -2.356 -18.035 -16.124 1.00 0. | 20 |
| MOTA | 2312 | | GLN | 157 | -1.447 -18.277 -15.847 1.00 0. | 20 |
| ATOM | 2313 | CA | GLN | 157 | -3.498 -18.941 -15.824 1.00 0. | 22 |
| ATOM | 2314 | HA | GLN | 157 | -3.987 -19.214 -16.747 1.00 0. | 24 |
| ATOM | 2315 | CB | GLN | 157 | -2.995 -20.204 -15.117 1.00 0. | |
| ATOM | 2316 | HB1 | | 157 | | 26 |
| ATOM | 2317 | _ | | 157 | A A 4 A A A A A A A A A A A A A A A A A | |
| ATOM | 2318 | | GLN | 157 | | 23 |
| ATOM | | | | | | 25 |
| | 2319 | | | 157 | | 94 |
| ATOM | 2320 | _ | | 157 | | 87 |
| MOTA | 2321 | | GLN | 157 | -2.152 -22.510 -15.598 1.00 1. | 19 |
| atom | 2322 | | GLN | 157 | -2.594 -22.799 -14.504 1.00 1. | 89 |
| MOTA | 2323 | NE2 | GLN | 157 | | 96 |
| MOTA | 2324 | HE21 | GLN | 157 | | 18 |
| MOTA | 2325 | | | 157 | | 65 |
| ATOM | 2326 | - | GLN | 157 | | |
| ATOM | 2327 | | GLN | 157 | · · - · - · - · - · - · - · - · | 22 |
| ATOM | 2328 | | | | | 24 |
| | | | GLY | 158 | -4.027 -17.456 -13.974 1.00 0. | - |
| MOTA | 2329 | | GLY | 158 | | 20 |
| ATOM | 2330 | | GLY | 158 | | 22 |
| MOTA | 2331 | | GLY | 158 | -4.380 -16.319 -12.232 1.00 0. | 22 |
| MOTA | 2332 | HA2 | GLY | 158 | | 25 |
| MOTA | 2333 | C | GLY | 158 | | 20 |
| ATOM | 2334 | 0 | GLY | 158 | • | 21 |
| ATOM | 2335 | | ILE | 159 | | - |
| ATOM | 2336 | | | | | 18 |
| | | - | ILE | 159 | | 18 |
| MOTA | 2337 | | ILE | 159 | | 19 |
| ATOM | 2338 | | ILE | 159 | | 20 |
| MOTA | 2339 | _ | ILE | 159 | | 19 |
| ATOM | 2340 | | ILE | 159 | | 20 |
| ATOM | 2341 | | | 159 | | 24 |
| MOTA | | HG11 | | 159 | | |
| ATOM | | HG12 | | 159 | | 26 |
| ATOM | 2344 | | ILE | | | 28 |
| ATOM | | _ | | 159 150 | | 21 |
| | | HG21 | | 159 | | 01 |
| ATOM | | HG22 | | 159 | -4.556 -13.274 -17.781 1.00 1. | 01 |
| ATOM | | HG23 | | 159 | -3.848 -14.398 -16.628 1.00 1. | 04 |
| ATOM | 2348 | | ILE | 159 | -5.571 -10.356 -15:166 1.00 O. | 27 |
| ATOM | 2349 | HD11 | .ILE | 159 | 4 | 05 |

| | | | | 450 | |
|-------|--------|--------|-------|------|---|
| MOTA | 2350 F | ID12 : | ILE | 159 | -4.644 -9.838 -14.978 1.00 1.06 |
| ATOM | 2351 B | ID13 : | ILE | 159 | -5.893 -10.848 -14.265 1.00 1.02 |
| MOTA | 2352 | | ILE | 159 | -6.644 -14.162 -16.173 1.00 0.21 |
| | | | | | |
| MOTA | 2353 | | ILE | 159 | -7.754 -13.700 -16.347 1.00 0.23 |
| MOTA | 2354 | N (| GLN | 160 | -6.215 -15.168 -16.885 1.00 0.22 |
| MOTA | 2355 | HN (| GLN | 160 | -5.322 -15.538 -16.726 1.00 0.21 |
| | | | | 160 | |
| MOTA | 2356 | | GLN | | -7.097 -15.763 -17.930 1.00 0.27 |
| MOTA | 2357 | HA (| GLN | 160 | -7.457 -14.979 -18.580 1.00 0.29 |
| MOTA | 2358 | CB (| GLN | 160 | -6.317 -16.786 -18.756 1.00 0.31 |
| | _ | | | | |
| MOTA | 2359 | | GLN | 160 | -6.999 -17.334 -19.389 1.00 0.35 |
| ATOM | 2360 | HB2 | GLN | 160 | -5.809 -17.472 -18.093 1.00 0.30 |
| ATOM | 2361 | CG | GLN | 160 | -5.289 -16.062 -19.626 1.00 0.34 |
| | | | | | |
| MOTA | 2362 | | GLN | 160 | -4.606 -15.512 -18.997 1.00 0.92 |
| ATOM | 2363 | HG2 | GLN | 160 | -5.799 -15.378 -20.290 1.00 0.91 |
| ATOM | 2364 | CD | GLN | 160 | -4.508 -17.087 -20.451 1.00 1.11 |
| | | | | | |
| MOTA | 2365 | | GLN | 160 | -4.451 -18.248 -20.100 1.00 1.88 |
| MOTA | 2366 | NE2 | GLN | 160 | -3.901 -16.704 -21.540 1.00 1.83 |
| ATOM | 2367 | HE21 | GLN | 160 | -3.947 -15.767 -21.824 1.00 2.13 |
| | | | | | |
| MOTA | | | GLN | 160 | -3.398 -17.353 -22.075 1.00 2.46 |
| MOTA | 2369 | С | GLN | 160 | -8.290 -16.447 -17.261 1.00 0.28 |
| MOTA | 2370 | 0 | GLN | .160 | -9.386 -16.449 -17.779 1.00 0.31 |
| | 2371 | - | | 161 | |
| MOTA | | N | SER | | -8.086 -17.035 -16.117 1.00 0.27 |
| MOTA | 2372 | HN | SER | 161 | -7.193 -17.030 -15.714 1.00 0.25 |
| ATOM | 2373 | CA | SER | 161 | -9.213 -17.718 -15.424 1.00 0.30 |
| ATOM | 2374 | | SER | 161 | |
| | | | | | -9.658 -18.444 -16.089 1.00 0.34 |
| ATOM | 2375 | CB | SER | 161 | -8.690 -18.427 -14.174 1.00 0.33 |
| ATOM | 2376 | HB1 | SER | 161 | -7.861 -19.067 -14.444 1.00 0.35 |
| | | | | | |
| MOTA | 2377 | | SER | 161 | -9.476 -19.024 -13.741 1.00 0.36 |
| MOTA | 2378 | OG | SER | 161 | -8.267 -17.455 -13.227 1.00 0.33 |
| ATOM | 2379 | HG | SER | 161 | -9.045 -16.986 -12.915 1.00 0.94 |
| ATOM | 2380 | C | SER | 161 | |
| | | | | | -10.267 -16.684 -15.019 1.00 0.30 |
| MOTA | 2381 | 0 | SER | 161 | -11.433 -16.997 -14.882 1.00 0.35 |
| ATOM | 2382 | N | LEU | 162 | -9.867 -15.457 -14.815 1.00 0.27 |
| MOTA | 2383 | HN | LEU | 162 | |
| | | | - | | -8.920 -15.225 -14.921 1.00 0.26 |
| ATOM | 2384 | CA | LEU | 162 | -10.852 -14.413 -14.405 1.00 0.29 |
| ATOM | 2385 | HA | LEU | 162 | -11.637 -14.869 -13.821 1.00 0.33 |
| ATOM | 2386 | CB | LEU | 162 | |
| | | | | | -10.141 -13.350 -13.563 1.00 0.28 |
| ATOM | 2387 | HB1 | LEU | 162 | -10.802 -12.509 -13.411 1.00 0.29 |
| ATOM | 2388 | HB2 | LEU | 162 | -9.256 -13.017 -14.086 1.00 0.27 |
| | | | | | |
| MOTA | 2389 | CG | LEU | 162 | -9.736 -13.937 -12.206 1.00 0.30 |
| MOTA | 2390 | HG | LEU | 162 | -9.157 -14.836 -12.367 1.00 0.30 |
| MOTA | 2391 | CD1 | LEU | 162 | -8.883 -12.918 -11.450 1.00 0.33 |
| ATOM | | HD11 | | 162 | |
| | | | | | -8.496 -13.370 -10.549 1.00 1.03 |
| MOTA | 2393 | HD12 | LEU | 162 | -9.490 -12.063 -11.191 1.00 1.01 |
| MOTA | 2394 | HD13 | LEU | 162 | -8.062 -12.601 -12.075 1.00 1.12 |
| ATOM | 2395 | CD2 | _ | | |
| | | | | 162 | -10.980 -14.272 -11.374 1.00 0.33 |
| MOTA | 2396 | HD21 | LEU | 162 | -11.227 -15.315 -11.502 1.00 1.05 |
| MOTA | 2397 | HD22 | LEU | 162 | -11.812 -13.664 -11.697 1.00 1.09 |
| MOTA | 2398 | HD23 | | 162 | |
| - | | | | | -10.776 -14.078 -10.332 1.00 1.01 |
| MOTA | 2399 | C | LEU | 162 | -11.461 -13.742 -15.643 1.00 0.30 |
| ATOM | 2400 | 0 | LEU | 162 | -12.664 -13.615 -15.757 1.00 0.36 |
| MOTA | 2401 | N | TYR | 163 | -10.645 -13.300 -16.564 1.00 0.27 |
| | | | | | |
| MOTA | 2402 | HN | TYR | 163 | -9.677 -13.404 -16.452 1.00 0.26 |
| MOTA | 2403 | CA | TYR | 163 | -11.188 -12.626 -17.783 1.00 0.31 |
| ATOM | 2404 | HA | TYR | 163 | -12.144 -12.182 -17.549 1.00 0.33 |
| ATOM | | СВ | | | |
| | 2405 | | TYR | 163 | -10.219 -11.531 -18.236 1.00 0.29 |
| MOTA | 2406 | HB1 | TYR | 163 | -10.562 -11.112 -19.170 1.00 0.32 |
| ATOM | 2407 | HB2 | TYR | 163 | -9.234 -11.952 -18.371 1.00 0.29 |
| MOTA | 2408 | CG | TYR | 163 | |
| | | | | | -10.162 -10.444 -17.190 1.00 0.25 |
| MOTA | 2409 | CD1 | TYR | 163 | -9.223 -10.520 -16.155 1.00 0.23 |
| ATOM | 2410 | HD1 | TYR | 163 | -8.545 -11.359 -16.103 1.00 0.23 |
| ATOM | 2411 | | TYR | 163 | |
| | | | | | |
| MOTA | 2412 | HD2 | TYR | 163 | -11.767 -9.298 -18.056 1.00 0.30 |
| ATOM | 2413 | CE1 | TYR | 163 | -9.164 -9.511 -15.187 1.00 0.24 |
| ATOM | 2414 | | TYR | 163 | |
| | | | | | |
| MOTA | 2415 | | TYR | 163 | -10.984 -8.348 -16.289 1.00 0.27 |
| MOTA | 2416 | HE2 | TYR | 163 | -11.663 -7.510 -16.340 1.00 0.30 |
| ATOM | 2417 | CZ | TYR | 163 | |
| | | | | | |
| MOTA | 2418 | OH | TYR | 163 | -9.985 -7.430 -14.299 1.00 0.31 |
| ATOM | 2419 | HH | TYR | 163 | -10.344 -7.782 -13.481 1.00 0.99 |
| ATOM | 2420 | C | TYR | 163 | |
| | | | | | |
| MOTA | 2421 | 0 | TYR | 163 | -11.953 -13.357 -19.933 1.00 0.43 |
| MOTA | 2422 | N | GLY | 164 | -10.865 -14.836 -18.729 1.00 0.38 |
| ATOM | 2423 | HN | GLY | 164 | |
| | | | | | |
| MOTA | 2424 | CA | GLY | 164 | -11.001 -15.877 -19.789 1.00 0.47 |
| MOTA | 2425 | HA1 | GLY | 164 | -11.851 -15.651 -20.413 1.00 0.53 |
| MOVIE | 2425 | | 01.14 | 164 | 11 140 16 044 10 006 100 0 0 |

| MOTA | 2427 | C | GLY | 164 | -9.735 | -15.902 | -20.648 | 1.00 | 0.55 |
|--------|-------|-------------|-----|-----|--------|---------|-----------------------|------|--|
| MOTA | 2428 | 0 | GLY | 164 | | | -21.819 | 1.00 | 1.01 |
| TER | 2429 | | GLY | 164 | | | | | |
| HETATM | | ZN | ZN | 166 | -0.218 | -6.515 | -2.613 | 1.00 | 0.24 |
| HETATM | | ZN | ZN | 167 | -3.506 | 6.833 | -0.714 | 1.00 | 0.24 |
| HETATM | | CA | CA | 168 | _ | | | | |
| HETATM | | Č1 | WAY | 169 | 6.060 | 3.350 | 3.030 | 1.00 | 0.23 |
| HETATM | | | , | | 2.180 | -4.315 | | 0.00 | 0.30 |
| | | C2 | WAY | 169 | 0.865 | -4.629 | | 0.00 | 0.33 |
| HETATM | | 1CE1 | | 169 | -0.170 | -4.517 | 2.143 | 0.00 | 0.38 |
| HETATM | | | YAW | 169 | 0.074 | -4.157 | 3.457 | 0.00 | 0.40 |
| HETATM | | 1CE2 | | 169 | 1.355 | -3.807 | 3.841 | 0.00 | 0.38 |
| HETATM | | C6 | WAY | 169 | 2.395 | -3.805 | 2.922 | 0.00 | 0.33 |
| HETATM | 2439 | 1HE1 | WAY | 169 | -1.190 | -4.713 | 1.839 | 0.00 | 0.42 |
| HETATM | 2440 | 1HZ | WAY | 169 | -0.734 | -4.151 | | 0.00 | 0.45 |
| HETATM | 2441 | 1HE2 | WAY | 169 | 1.535 | -3.534 | | 0.00 | 0.42 |
| HETATM | 2442 | C10 | WAY | 169 | 0.444 | -5.080 | -0.136 | 0.00 | 0.36 |
| HETATM | 2443 | | WAY | 169 | 0.467 | -6.264 | | 0.00 | 0.58 |
| HETATM | | | WAY | 169 | -0.019 | -4.195 | _ | | |
| HETATM | | | WAY | 169 | -0.045 | | | 0.00 | 0.61 |
| HETATM | | | WAY | 169 | | -4.608 | -2.371 | 0.00 | 0.68 |
| HETATM | | | | | -0.357 | -3.297 | -0.743 | 0.00 | 0.88 |
| | | | WAY | 169 | -0.953 | -4.727 | | 0.00 | 1.13 |
| HETATM | | 1CH1 | | 169 | 3.728 | -3.247 | 3.360 | 0.00 | 0.37 |
| HETATM | | 1HH1 | | 169 | 3.702 | -2.162 | 3.422 | 0.00 | 1.07 |
| HETATM | . – – | 1HH2 | | 169 | 4.519 | -3.516 | 2.664 | 0.00 | 1.06 |
| HETATM | | 1HH3 | | 169 | 4.013 | -3.623 | 4.339 | 0.00 | 1.11 |
| HETATM | 2452 | N20 | WAY | 169 | 3.274 | -4.485 | 0.819 | 0.00 | 0.29 |
| HETATM | 2453 | S21 | WAY | 169 | 3.865 | -3.175 | - | 0.00 | 0.25 |
| HETATM | 2454 | 2CB | WAY | 169 | 3.882 | -5.812 | 0.684 | 0.00 | 0.32 |
| HETATM | 2455 | 2CE1 | | 169 | 7.334 | -6.241 | 2.178 | 0.00 | 1.09 |
| HETATM | | | | | 6.971 | -6.520 | | | and the second s |
| HETATM | | | WAY | 169 | | | | 0.00 | 0.53 |
| HETATM | | | | 169 | 5.697 | -6.659 | | 0.00 | |
| HETATM | | | | | 4.747 | -6.451 | | 0.00 | |
| | | | | 169 | 5.010 | -6.084 | | 0.00 | |
| HETATM | | | | 169 | 6.338 | -5.982 | | 0.00 | 1.14 |
| HETATM | | | | 169 | 8.374 | -6.224 | | 0.00 | 1.94 |
| HETATM | | | WAY | 169 | 7.752 | -6.630 | 4.227 | 0.00 | 0.61 |
| HETATM | | | | 169 | 3.708 | -6.570 | 3.227 | 0.00 | 2.23 |
| HETATM | | | | 169 | 6.599 | -5.706 | 0.239 | 0.00 | _ |
| HETATM | 2465 | 2HB1 | WAY | 169 | 4.245 | -5.905 | | 0.00 | |
| HETATM | 2466 | 2HB2 | WAY | 169 | 3.095 | -6.552 | | 0.00 | 0.34 |
| HETATM | | | | 169 | 4.187 | -3.617 | | 0.00 | |
| HETATM | | | | 169 | 3.310 | -3.216 | | 0.00 | |
| HETATM | | | | 169 | 3.622 | | | 0.00 | |
| HETATM | | | | 169 | 4.769 | -4.183 | | | 0:27 |
| HETATM | | | | 169 | | | | 0.00 | 0.24 |
| HETATM | | | | | 5.602 | -4.644 | | 0.00 | |
| | | | | 169 | 5.315 | -4.359 | ·- · - · - | 0.00 | 0.23 |
| HETATM | | | | 169 | 2.392 | -2.714 | + + | 0.00 | 0.29 |
| HETATM | | | | 169 | 2.961 | -3.091 | | 0.00 | 0.31 |
| HETATM | | | | 169 | 6.481 | -5.228 | -3.535 | 0.00 | 0.26 |
| HETATM | | | | 169 | 5.959 | -4.707 | -1.184 | 0.00 | 0.27 |
| HETATM | | | WAY | 169 | 5.078 | -4.439 | -5.664 | 0.00 | 0.27 |
| HETATM | | | | 169 | 6.245 | -5.202 | | 0.00 | 0.28 |
| HETATM | | | | 169 | 6.379 | | | 0.00 | 0.31 |
| HETATM | 2480 | 3HH2 | WAY | 169 | 6.178 | -6.172 | | 0.00 | 0.28 |
| HETATM | | | | 169 | 7.127 | | | 0.00 | |
| HETATM | | | WAY | 169 | 5.123 | -2.847 | | 0.00 | 0.23 |
| HETATM | | | WAY | 169 | 2.834 | -2.186 | | 0.00 | |
| END | | ~ ~ | | | 2.034 | 2,100 | 0.004 | 0.00 | 0.25 |
| | | | | | | | | | |

| | A | tom | Res | • | x | Y | z | Occ. B | MOT |
|--------------|----------|-----------|------------|----------|-------------------|------------------|------------------|---------------------------|----------------|
| ATOM | | ype CB | THR | 7 | 73.468 | 27.410 | 6.079 | Occ. B | MOL. A_13 |
| MOTA | 2 | OG1 | THR | 7 | 72.149 | 27.911 | 6.358 | 1.00 37.82 | A_13 |
| ATOM ATOM | | | THR THR | 7 7 | 73.843 75.936 | 26.297 28.076 | 7.068 6.227 | 1.00 25.79 1.00 28.29 | A_13 |
| ATOM | | | THR | ż | 76.497 | 28.070 | 7.332 | 1.00 28.29 .1.00 22.94 | A_13 A_13 |
| ATOM | 9 | N | THR | 7 | 74.360 | 29.396 | 4.862 | 1.00 20.25 | A_13 |
| ATOM ATOM | | | THR LEU | 7 8 | 74.501 76.547 | 28.593 | 6.099 | 1.00 21.49 | A_13 |
| ATOM | 14 | | LEU | 8 | 77.915 | 27.691 27.150 | 5.099 5.105 | 1.00 32.90 1.00 31.85 | · A_13 A_13 |
| MOTA | | CB | LEU | 8 | 77.952 | 25.759 | 4.438 | 1.00 21.38 | A_13 |
| ATOM ATOM | 16 17 | | LEU LEU | 8 8 | 78.016 79.463 | 25.576 | 2.910 | 1.00 29.31 | A_13 |
| ATOM | 18 | | LEU | 8 | 77.334 | 25.509 24.292 | 2.425 2.527 | 1.00 16.78 1.00 23.37 | A_13 A_13 |
| MOTA | 19 | C | LEU | 8 | 78.956 | 28.070 | 4.465 | 1.00 24.01 | A_13 |
| MOTA MOTA | 20 21 | O N | LEU LYS | 8 9 | 78.835 | 28.415 | 3.293 | 1.00 26.18 | A_13 |
| ATOM | 23 | CA | LYS | 9 | 79.980 81.106 | 28.424 29.298 | 5.251 4.867 | 1.00 36.26 1.00 23.24 | A_13 A_13 |
| ATOM | 24 | CB | LYS | 9 | 82.438 | 28.521 | 4.977 | 1.00 25.52 | A_13 |
| ATOM | 25 26 | CG | LYS | 9 9 | 82.767 | 27.570 | 3.815 | 1.00 19.05 | A_13 |
| MOTA MOTA | 27 | CE | LYS LYS | 9 | 83.661 83.451 | 28.243 27.688 | 2.753 1.323 | 1.00 31.69 1.00 25.30 | A_13 A_13 |
| MOTA | 28 | NZ | LYS | 9 | 82.056 | 27.938 | 0.797 | 1.00 20.65 | A_13 |
| ATOM ATOM | 32 33 | 0 | LYS LYS | 9 9 | 81.042 | 30.073 | 3.526 | 1.00 31.41 | A_13 |
| MOTA | 34 | N | TRP | 10 | 80.764 81.327 | 29.505 31.372 | 2.466 3.573 | 1.00 22.31 1.00 15.84 | A_13 A_13 |
| ATOM | 36 | CA | TRP | 10 | 81.312 | 32.172 | 2.361 | 1.00 10.58 | A_13 |
| ATOM ATOM | 37 38 | CB | TRP TRP | 10 10 | 81.636 80.529 | 33.620 | 2.680 | 1.00 21.39 | A_13 |
| ATOM | 39 | CD2 | TRP | 10 | 79.479 | 34.337 35.074 | 3.343 2.697 | 1.00 22.84 1.00 20.41 | A_13 A_13 |
| MOTA | 40 | CE2 | TRP | 10 | 78.676 | 35.631 | 3.718 | 1.00 24.50 | A_13 |
| ATOM ATOM | 41 42 | CE3 | TRP | 10 10 | 79.142 | 35.320 | 1.357 | 1.00 13.29 | A_13 |
| ATOM | 43 | NE1 | | 10 | 80.327 79.220 | 34.469 35.253 | 4.682 4.919 | 1.00 13.40 1.00 18.40 | A_13 A_13 |
| MOTA | 45 | CZ2 | TRP | 10 | 77.550 | 36.418 | 3.442 | 1.00 12.63 | A_13 |
| MOTA | 46 47 | CZ3 | | 10 | 78.021 | 36.105 | 1.083 | 1.00 19.89 | A_13 |
| MOTA MOTA | 48 | CH2 C | TRP TRP | 10 10 | 77.242 82.377 | 36.641 31.594 | 2.120 1.455 | 1.00 13.62 1.00 22.95 | A_13 A_13 |
| MOTA, | 49 | 0 | TRP | 10 | 83.450 | 31.221 | 1.920 | 1.00 16.28 | A_13 |
| ATOM | 50 52 | N | SER | 11 | 82.087 | 31.533 | 0.167 | 1.00 14.81 | A_13 |
| MOTA MOTA | 53 | CA | SER SER | 11 11 | 83.017 82.282 | 30.975 30.596 | -0.801 -2.086 | 1.00 19.50 1.00 24.36 | A_13 A_13 |
| MOTA | 54 | OG | SER | 11 | 81.605 | 29.353 | -1.958 | 1.00 40.49 | A_13 |
| MOTA MOTA | 56 57 | С О | SER | 11 | 84.190 | 31.867 | -1.134 | 1.00 16.53 | A_13 |
| ATOM | 58 | N | ser Lys | 11 12 | 85.132 84.153 | 31.423 33.113 | -1.779 -0.686 | 1.00 23.48 1.00 12.50 | A_13 A_13 |
| MOTA | 60 | CA | LYS | 12 | 85.232 | 34.057 | -0.961 | 1.00 17.05 | A_13 |
| MOTA MOTA | 61 62 | CB CG | LYS LYS | 12 | 84.741 | 35.168 | -1.891 | 1.00 17.32 | A_13 |
| ATOM | 63 | CD | LYS | 12 12 | 83.526 82.788 | 35.898 36.644 | -1.350 -2.446 | 1.00 18.49 1.00 18.29 | A_13 A_13 |
| MOTA | 64 | CE | LYS | 12 | 81.534 | 37.282 | -1.888 | 1.00 18.44 | A_13 |
| MOTA MOTA | 65 69 | NZ | LYS LYS | 12 12 | 80.805 | 38.094 | -2.895 | 1.00 16.65 | A_13 |
| ATOM | 70 | Ö | LYS | 12 | 85.687 84.946 | 34.662 34.637 | 0.344 1.319 | 1.00 11.16 1.00 12.63 | A_13 A_13 |
| MOTA | 71 | N | MET | 13 | 85.915 | 35.185 | 0.355 | 1.00 15.52 | A_13. |
| MOTA MOTA | 73 74 | CA CB | MET | 13 13 | 87.516. 89.028 | 35.801 35.547 | 1.537 1.565 | 1.00 11.04 1.00 16.57 | A_13 |
| MOTA | 75 | CG | MET | 13 | 89.431 | 34.082 | 1.707 | 1.00 20.92 | A_13 A_13 |
| MOTA | 76 | SD | MET | 13 | 88.905 | 33.235 | 3.227 | 1.00 20.10 | A_13 |
| MOTA MOTA | 77 78 | CE | MET | 13 13 | 87.486 87.258 | 32.313 37.296 | 2.604 1.572 | 1.00 16.29 1.00 13.23 | A_13 A_13 |
| MOTA | 79 | ŏ | MET | 13 | 87.247 | 37.916 | 2.634 | 1.00 22.80 | A_13 |
| MOTA | 80 | N | ASN | 14 | 87.111 | 37.875 | 0.389 | 1.00 15.02 | A_13 |
| MOTA MOTA | 82 83 | CA | asn asn | 14 14 | 86.853 87.445 | 39.294 39.801 | 0.241 -1.082 | 1.00 33.02 1.00 19.42 | A_13 |
| MOTA | 84 | CG | ASN | 14 | 88.925 | 39.482 | -1.217 | 1.00 30.32 | A_13 A_13 |
| MOTA | 85 | OD1 | ASN | 14 | 89.343 | 38.341 | -1.031 | 1.00 30.12 | A_13 |
| MOTA MOTA | 86 89 | ND2 C | ASN ASN | 14 | 89.723 85.337 | 40.489 | -1.549 | 1.00 28.22 | A_13 |
| ATOM | 90 | 0 | asn | 14 14 | 84.606 | 39.482 38.935 | 0.277 -0.568 | 1.00 27.58 1.00 28.01 | A_13 A_13 |
| MOTA | 91 | N | LEU | 15 | 84.868 | 40.212 | 1.287 | 1.00 19.06 | A_13 |
| MOTA MOTA | 93 94 | CA CB | LEU | 15 15 | 83.444 | 40.450 | 1.459 | | A_13 |
| ATOM | 95 | CG | Leu Leu | 15 15 | 82.930 83.027 | 39.690 38.166 | 2.691 2.593 | | A_13 A_13 |
| MOTA | 96 | CD1 | LEU | 15 | 83.216 | 37.555 | 3.962 | 1.00 17.48 | A_13 |
| MOTA MOTA | 97 98 | CD2 C | LEU | 15 15 | 81.799 83.161 | | 1.903 | | A_13 |
| ATOM | 99 | Ö | LEU | 15 | 83.980 | | 1.609 2.130 | | A_13 A_13 |

FIG. 5

| 7 TOM | 100 | | m.15 | | 01 000 | | | | |
|-------|-----|-----|------|----|--------|-------------|--------|------------|------|
| ATOM | 100 | N | THR | 16 | 81.983 | 42.343 | 1.162 | 1.00 21.22 | A_13 |
| ATOM | 102 | CA | THR | 16 | 81.578 | 43.736 | 1.252 | 1.00 10.00 | A_13 |
| ATOM | 103 | CB | THR | 16 | 81.194 | 44.257 | -0.109 | 1.00 10.00 | A_13 |
| MOTA | 104 | OG1 | THR | 16 | 80.225 | 43.370 | -0.681 | 1.00 22.43 | A_13 |
| ATOM | 106 | CG2 | THR | 16 | 82.427 | 44.383 | -1.009 | 1.00 15.42 | A_13 |
| ATOM | 107 | C | THR | 16 | 80.368 | 43.869 | 2.184 | 1.00 14.48 | A_13 |
| MOTA | 108 | 0 | THR | 16 | 79.647 | 42.897 | 2.445 | 1.00 15.74 | A_13 |
| MOTA | 109 | N | TYR | 17 | 80.176 | 45.065 | 2.716 | 1.00 15.89 | A_13 |
| ATOM | 111 | CA | TYR | 17 | 79.064 | 45.340 | 3.604 | 1.00 13.19 | |
| ATOM | 112 | CB | TYR | 17 | 79.480 | 45.195 | | | A_13 |
| MOTA | 113 | CG | TYR | 17 | 80.448 | | 5.067 | 1.00 21.42 | A_13 |
| ATOM | 114 | CD1 | | | | 46.236 | 5.580 | 1.00 26.23 | A_13 |
| | | | TYR | 17 | 81.824 | 46.081 | 5.412 | 1.00 16.37 | A_13 |
| ATOM | 115 | CE1 | TYR | 17 | 82.724 | 46.981 | 5.988 | 1.00 12.90 | A_13 |
| ATOM | 116 | CD2 | TYR | 17 | 79.990 | 47.329 | 6.331 | 1.00 17.15 | A_13 |
| MOTA | 117 | CE2 | TYR | 17 | 80.880 | 48.235 | 6.912 | 1.00 24.15 | A_13 |
| MOTA | 118 | CZ | TYR | 17 | 82.244 | 48.057 | 6.743 | 1.00 23.38 | A_13 |
| ATOM | 119 | OH | TYR | 17 | 83.121 | 48.942 | 7.343 | 1.00 19.47 | A_13 |
| MOTA | 121 | C | TYR | 17 | 78.573 | 46.740 | 3.343 | 1.00 10.00 | A_13 |
| ATOM | 122 | 0 | TYR | 17 | 79.298 | 47.559 | 2.782 | 1.00 19.27 | A_13 |
| ATOM | 123 | N | ARG | 18 | 77.349 | 47.019 | 3.762 | 1.00 18.52 | |
| MOTA | 125 | CA | ARG | 18 | 76.762 | 48.332 | 3.577 | 1.00 10.00 | A_13 |
| MOTA | 126 | CB | ARG | 18 | 75.970 | 48.363 | 2.274 | | A_13 |
| ATOM | 127 | CG | ARG | 18 | 75.134 | | | 1.00 10.00 | A_13 |
| ATOM | 128 | CD | ARG | | | 49.619 | 2.094 | 1.00 14.01 | A_13 |
| ATOM | 129 | | | 18 | 74.266 | 49.524 | 0.846 | 1.00 13.91 | A_13 |
| | | NE | ARG | 18 | 73.298 | 50.615 | 0.782 | 1.00 13.55 | A_13 |
| MOTA | 131 | CZ | ARG | 18 | 72.165 | 50.571 | 0.092 | 1.00 10.00 | A_13 |
| ATOM | 132 | NH1 | ARG | 18 | 71.855 | 49.488 | -0.602 | 1.00 14.30 | A_13 |
| MOTA | 135 | NH2 | ARG | 18 | 71.331 | 51.604 | 0.125 | 1.00 28.79 | A_13 |
| ATOM | 138 | C | ARG | 18 | 75.842 | 48.640 | 4.741 | 1.00 10.65 | A_13 |
| ATOM | 139 | Ο. | ARG | 18 | 75.037 | 47.796 | 5.141 | 1.00 12.86 | A_13 |
| ATOM | 140 | N | ILE | 19 | 76.014 | 49.814 | 5.332 | 1.00 25.54 | A_13 |
| ATOM | 142 | CA | ILE | 19 | 75.169 | 50.265 | 6.436 | 1.00 24.52 | A_13 |
| ATOM | 143 | CB | ILE | 19 | 75.944 | | 7.350 | 1.00 18.37 | |
| ATOM | 144 | | ILE | 19 | 75.034 | 51.765 | 8.485 | | A_13 |
| ATOM | 145 | | ILE | 19 | 77.204 | | | 1.00 13.87 | A_13 |
| ATOM | 146 | CD1 | ILE | | | 50.545 | 7.888 | 1.00 27.67 | A_13 |
| ATOM | 147 | | | 19 | 78.203 | 51.501 | 8.557 | 1.00 22.81 | A_13 |
| | | C | ILE | 19 | 74.062 | | 5.698 | 1.00 21.11 | A_13 |
| ATOM | 148 | 0 | ILE | 19 | 74.261 | 52.179 | 5.300 | 1.00 10.00 | A_13 |
| MOTA | 149 | N | LAV | 20 | 72.916 | 50.378 | 5.487 | 1.00 19.76 | A_13 |
| MOTA | 151 | CA | VAL | 20 | 71.829 | 51.014 | 4.735 | 1.00 18.20 | A_13 |
| ATOM | 152 | CB | VAL | 20 | 70.774 | 49.983 | 4.193 | 1.00 15.42 | A_13 |
| MOTA | 153 | CG1 | VAL | 20 | 71.384 | 48.570 | 4.088 | 1.00 10.00 | A_13 |
| ATOM | 154 | | VAL | 20 | 69.496 | 50.030 | 4.992 | 1.00 18.62 | |
| ATOM | 155 | C | VAL | 20 | 71.175 | | 5.443 | 1.00 18.62 | A_13 |
| ATOM | 156 | O | VAL | 20 | 70.652 | 53.110 | 4.798 | | A_13 |
| ATOM | 157 | N | ASN | 21 | 71.153 | 52.187 | | 1.00 18.36 | A_13 |
| ATOM | 159 | CA | ASN | 21 | | | 6.773 | 1.00 10.94 | A_13 |
| ATOM | 160 | CB | | | 70.609 | 53.316 | 7.544 | 1.00 11.99 | A_13 |
| ATOM | | | ASN | 21 | 69.078 | 53.307 | 7.675 | 1.00 10.00 | A_13 |
| | 161 | CG | ASN | 21 | 68.533 | 51.978 | 8.107 | 1.00 14.93 | A_13 |
| ATOM | 162 | | ASN | 21 | 67.627 | 51.449 | 7.486 | 1.00 21.54 | A_13 |
| ATOM | 163 | | ASN | 21 | 69.105 | · · · · · · | 9.148 | 1.00 10.00 | A_13 |
| ATOM | 166 | C | ASN | 21 | 71.291 | 53.382 | 8.897 | 1.00 18.90 | A_13 |
| ATOM | 167 | 0 | ASN | 21 | 72.006 | 52.447 | 9.283 | 1.00 12.49 | A_13 |
| MOTA | 168 | N | TYR | 22 | 71.053 | 54.471 | 9.618 | 1.00 17.47 | A_13 |
| MOTA | 170 | CA | TYR | 22 | 71.681 | 54.708 | 10.910 | 1.00 24.85 | A_13 |
| ATOM | 171 | CB | TYR | 22 | 72.556 | 55.954 | 10.818 | 1.00 13.52 | A_13 |
| MOTA | 172 | CG | TYR | 22 | 73.791 | 55.748 | 9.991 | 1.00 10.00 | A_13 |
| ATOM | 173 | CD1 | TYR | 22 | 75.033 | 55.600 | 10.598 | 1.00 14.05 | |
| ATOM | 174 | CE1 | | 22 | 76.180 | 55.370 | 9.841 | 1.00 13.69 | A_13 |
| ATOM | 175 | | TYR | 22 | 73.717 | 55.663 | | | A_13 |
| ATOM | 176 | CE2 | | 22 | 74.848 | | 8.608 | 1.00 10.00 | A_13 |
| ATOM | 177 | CZ | TYR | | | 55.432 | 7.847 | 1.00 17.10 | A_13 |
| ATOM | 178 | | | 22 | 76.077 | 55.288 | 8.476 | 1.00 14.43 | A_13 |
| | | OH | TYR | 22 | 77.204 | 55.072 | 7.737 | 1.00 10.00 | A_13 |
| ATOM | 180 | C | TYR | 22 | 70.726 | 54.862 | 12.076 | 1.00 25.95 | A_13 |
| ATOM | 181 | 0 | TYR | 22 | 69.593 | 55.311 | 11.916 | 1.00.10.00 | A_13 |
| ATOM | 182 | N | THR | 23 | 71.187 | 54.483 | 13.259 | 1.00 20.30 | A_13 |
| ATOM | 184 | CA | THR | 23 | 70.367 | 54.606 | 14.450 | 1.00 29.11 | A_13 |
| MOTA | 185 | CB | THR | 23 | 70.821 | 53.635 | 15.584 | 1.00 10.90 | A_13 |
| ATOM | 186 | OG1 | | 23 | 70.136 | 53.968 | 16.792 | 1.00 10.00 | |
| ATOM | 188 | CG2 | | 23 | 72.328 | 53.752 | 15.852 | | A_13 |
| ATOM | 189 | C | THR | 23 | 70.459 | 56.038 | | 1.00 16.51 | A_13 |
| ATOM | 190 | Ö | THR | 23 | 71.360 | | 14.959 | 1.00 18.14 | A_13 |
| ATOM | 191 | N | PRO | 24 | | 56.785 | 14.575 | 1.00 10.00 | A_13 |
| MOTA | 192 | CD | PRO | | 69.433 | 56.487 | 15.691 | 1.00 12.76 | A_13 |
| ATOM | 193 | | | 24 | 68.061 | 55.950 | 15.716 | 1.00 15.26 | A_13 |
| ATOM | | CA | PRO | 24 | 69.453 | 57.844 | 16.232 | 1.00 22.70 | A_13 |
| ATOM | 194 | CB | PRO | 24 | 67.985 | 58.086 | 16.585 | 1.00 28.52 | A_13 |
| ATOM | 195 | CG | PRO | 24 | 67.448 | 56.706 | 16.841 | 1.00 15.78 | A_13 |

| ATOM | 196 | С | PRO | 24 | 70.346 | 57.945 | 17.475 | 1.00 24.52 | A_13 |
|------|-----|-----|-----|----------|--------|--------|--------|------------|------|
| MOTA | 197 | 0 | PRO | 24 | 70.790 | 59.040 | 17.831 | 1.00 10.00 | A_13 |
| ATOM | 198 | N | ASP | 25 | 70.614 | 56.797 | 18.105 | 1.00 10.90 | |
| ATOM | 200 | CA' | ASP | 25 | | | | | A_13 |
| | | | | | 71.416 | 56.721 | 19.336 | 1.00 12.31 | A_13 |
| ATOM | 201 | CB | ASP | 25 | 71.339 | 55.317 | 19.917 | 1.00 25.26 | A_13 |
| ATOM | 202 | CG | ASP | 25 | 69.927 | 54.782 | 19.977 | 1.00 10.00 | A_13 |
| MOTA | 203 | OD1 | ASP | 25 | 69.783 | 53.567 | 20.159 | 1.00 20.90 | A_13 |
| MOTA | 204 | OD2 | ASP | 25 | 68.960 | 55.558 | 19.841 | 1.00 18.45 | A_13 |
| MOTA | 205 | С | ASP | 25 | 72.891 | 57.113 | 19.286 | 1.00 14.34 | A_13 |
| ATOM | 206 | Ö | ASP | 25 | 73.449 | 57.511 | 20.301 | 1.00 11.77 | |
| | 207 | | | | | | | | A_13 |
| MOTA | | N | MET | 26 26 | 73.546 | 56.873 | 18.157 | 1.00 20.78 | A_13 |
| MOTA | 209 | CA. | MET | 26 | 74.960 | 57.208 | 18.010 | 1.00 20.03 | A_13 |
| MOTA | 210 | CB | MET | 26 | 75.791 | 55.928 | 17.916 | 1.00 13.86 | A_13 |
| MOTA | 211 | CG | MET | 26 | 75.966 | 55.181 | 19.231 | 1.00 19.00 | A_13 |
| MOTA | 212 | SD | MET | 26 | 76.043 | 53.404 | 18.941 | 1.00 14.67 | A_13 |
| MOTA | 213 | CE | MET | 26 | 77.737 | 53.223 | 18.385 | 1.00 19.74 | A_13 |
| MOTA | 214 | C | MET | 26 | 75.157 | 58.047 | 16.754 | 1.00 13.32 | |
| ATOM | 215 | ŏ | MET | 26 | 74.274 | | | | A_13 |
| | | | | | | 58.086 | 15.900 | 1.00 16.81 | A_13 |
| ATOM | 216 | N | THR | 27 | 76.285 | 58.749 | 16.656 | 1.00 10.29 | A_13 |
| ATOM | 218 | CA | THR | 27 | 76.568 | 59.564 | 15.470 | 1.00 17.00 | A_13 |
| MOTA | 219 | CB | THR | 27 | 77.710 | 60.596 | 15.700 | 1.00 11.79 | A_13 |
| MOTA | 220 | OG1 | THR | 27 | 78.969 | 59.921 | 15.729 | 1.00 23.77 | A_13 |
| MOTA | 222 | CG2 | THR | 27 | 77.519 | 61.342 | 17.020 | 1.00 21.98 | A_13 |
| MOTA | 223 | C | THR | 27 | 76.996 | 58.634 | 14.347 | 1.00 13.37 | A_13 |
| MOTA | 224 | 0 | THR | 27 | 77.411 | 57.500 | 14.608 | 1.00 11.05 | |
| MOTA | 225 | N | HIS | 28 | 76.972 | 59.124 | 13.113 | 1.00 10.00 | A_13 |
| MOTA | 227 | CA | HIS | 28 | | | | | A_13 |
| ATOM | 228 | | | | 77.362 | 58.300 | 11.980 | 1.00 10.96 | A_13 |
| | | CB | HIS | 28 | 77.240 | 59.071 | 10.657 | 1.00 16.07 | A_13 |
| MOTA | 229 | CG | HIS | 28 | 75.829 | 59.382 | 10.264 | 1.00 15.53 | A_13 |
| MOTA | 230 | | HIS | 28 | 74.707 | 59.531 | 11.016 | 1.00 21.47 | A_13 |
| MOTA | 231 | ND1 | HIS | 28 | 75.440 | 59.597 | 8.959 | 1.00 30.32 | A_13 |
| ATOM | 233 | ÇE1 | HIS | 28 | 74.149 | 59.868 | 8.920 | 1.00 19.38 | A_13 |
| MOTA | 234 | NE2 | HIS | 28 | 73.680 | 59.833 | 10.160 | 1.00 29.43 | A_13 |
| ATOM | 236 | C | HIS | 28 | 78.769 | 57.735 | 12.151 | 1.00 14.80 | |
| ATOM | 237 | ŏ | HIS | 28 | 79.005 | | | | A_13 |
| ATOM | 238 | N | SER | | | 56.568 | | 1.00 28.24 | A_13 |
| | | | | 29 | 79.703 | 58.548 | 12.634 | 1.00 14.00 | A_13 |
| ATOM | 240 | CA | SER | 29 | 81.068 | 58.070 | 12.854 | 1.00 19.57 | A_13 |
| ATOM | 241 | CB | SER | 29 | 82.001 | 59.219 | 13.242 | 1.00.17.84 | A_13 |
| ATOM | 242 | OG | SER | 29 | 82.383 | 59.936 | 12.084 | 1.00 28.25 | A_13 |
| MOTA | 244 | C | SER | 29 | 81.134 | 56.983 | 13.917 | 1.00 15.23 | A_13 |
| MOTA | 245 | 0 | SER | 29 | 81.818 | 55.973 | 13.733 | 1.00 13.73 | A_13 |
| MOTA | 246 | N | GLU | 30 | 80.428 | 57.182 | 15.027 | 1.00 27.71 | - |
| ATOM | 248 | CA | GLU | 30 | 80.430 | 56.186 | | - | A_13 |
| ATOM | 249 | CB | GLU | 30 | | | 16.100 | 1.00 23.60 | A_13 |
| ATOM | 250 | | | | 79.571 | 56.635 | 17.289 | 1.00 21.72 | A_13 |
| | | CG | GLU | 30 | 80.048 | 57.913 | | 1.00 24.07 | A_13 |
| ATOM | 251 | CD | GLU | 30 | 79.205 | 58.279 | 19.185 | 1.00 21.06 | A_13 |
| MOTA | 252 | OE1 | | 30 | 79.784 | 58.660 | 20.218 | 1.00 46.95 | A_13 |
| MOTA | 253 | OE2 | GLU | 30 | 77.963 | 58.185 | 19.119 | 1.00 18.27 | A_13 |
| ATOM | 254 | C | GLU | 30 | 79.895 | 54.877 | | 1.00 18.75 | A_13 |
| MOTA | 255 | 0 | GLU | 30 | 80.456 | 53.809 | 15.815 | 1.00 13.06 | A_13 |
| ATOM | 256 | N | VAL | 31 | 78.839 | 54.970 | | 1.00 16.23 | |
| ATOM | 258 | CA | VAL | 31 | 78.225 | 53.781 | | | A_13 |
| MOTA | 259 | CB | VAL | 31 | 76.899 | | _ | | A_13 |
| ATOM | 260 | CG1 | | | | 54.135 | | 1.00 23.53 | A_13 |
| ATOM | 261 | CG2 | | 31 | 76.384 | 52.920 | 12.628 | 1.00 14.39 | A_13 |
| | | | | 31 | 75.829 | 54.587 | 14.377 | 1.00 10.00 | A_13 |
| MOTA | 262 | C | VAL | 31 | 79.208 | 53.040 | 13.216 | 1.00 20.29 | A_13 |
| MOTA | 263 | 0 | VAL | 31 | 79.330 | 51.814 | 13.282 | 1.00 14.02 | A_13 |
| MOTA | 264 | N | GLU | 32 | 79.913 | 53.790 | 12.370 | 1.00 23.94 | A_13 |
| ATOM | 266 | CA | GLU | 32 | 80.887 | 53.219 | 11.446 | 1.00 10.18 | A_13 |
| MOTA | 267 | CB | GLU | 32 | 81.406 | 54.285 | 10.502 | 1.00 16.50 | A_13 |
| MOTA | 268 | CG | GLU | 32 | 80.424 | | 9.427 | 1.00 20.84 | A_13 |
| MOTA | 269 | CD | GLU | 32 | 80.330 | 56.080 | 9.155 | 1.00 22.31 | |
| MOTA | 270 | OE1 | | 32 | 79.285 | | | | A_13 |
| ATOM | 271 | OE2 | | | | 56.509 | 8.639 | 1.00 29.39 | A_13 |
| | | | | 32 | 81.294 | 56.812 | 9.458 | 1.00 22.01 | A_13 |
| ATOM | 272 | C | GLU | 32 | 82.056 | 52.565 | | 1.00 18.93 | A_13 |
| ATOM | 273 | 0 | GLU | 32 | 82.474 | | 11.753 | 1.00 24.42 | A_13 |
| MOTA | 274 | N | LYS | 33 | 82.610 | 53.241 | 13.139 | 1.00 19.78 | A_13 |
| ATOM | 276 | CA | LYS | 33 | 83.726 | 52.661 | | 1.00 28.68 | A_13 |
| ATOM | 277 | CB | LYS | 33 | 84.340 | | | 1.00 18.54 | N 13 |
| MOTA | 278 | CG | LYS | 33 | 85.016 | 54.855 | | | A_13 |
| ATOM | 279 | CD | LYS | 33 | | | 14.135 | 1.00 31.19 | A_13 |
| ATOM | 280 | CE | LYS | | 86.135 | 54.425 | 13.148 | 1.00 40.31 | A_13 |
| MOTA | 281 | | | 33 | 85.600 | 53.972 | | | A_13 |
| | | NZ | LYS | 33 | 86.646 | 53.779 | | 1.00 33.20 | A_13 |
| ATOM | 285 | C | LYS | 33 | 83.242 | 51.407 | 14.594 | 1.00 12.66 | A_13 |
| ATOM | 286 | 0 | LYS | 33 | 83.892 | 50.361 | 14.552 | 1.00 15.54 | A_13 |
| ATOM | 287 | N | ALA | 34 | 82.036 | 51.481 | | 1.00 20.70 | A_13 |
| MOTA | 289 | CA | ALA | 34 | 81.453 | 50.344 | 15.843 | 1.00 10.00 | A_13 |
| | | | | | | | | | |

| MOTA | 290 | CB | ALA | 34 | 80.040 | 50.651 | 16.279 | 1.00 18.59 | A_13 |
|------|-----|-----|-----|----|--------|--------|--------|------------|--------|
| ATOM | 291 | С | ALA | 34 | 81.468 | 49.119 | 14.940 | 1.00 13.45 | |
| MOTA | 292 | 0 | ALA | 34 | 82.067 | 48.095 | | | A_13 |
| ATOM | 293 | N | | | | | 15.284 | 1.00 15.90 | A_13 |
| | | | PHE | 35 | 80.857 | 49.234 | 13.766 | 1.00 19.57 | A_13 |
| ATOM | 295 | CA | PHE | 35 | 80.802 | 48.112 | 12.812 | 1.00 26.77 | A_13 |
| MOTA | 296 | CB | PHE | 35 | 79.837 | 48.423 | 11.660 | 1.00 17.34 | A_13 |
| ATOM | 297 | CG | PHE | 35 | 78.390 | 48.477 | 12.077 | 1.00 30.55 | |
| MOTA | 298 | | PHE | 35 | | | | | A_13 |
| | | | | | 77.838 | 47.464 | 12.863 | 1.00 26.58 | A_13 |
| ATOM | 299 | | PHE | 35 | 77.570 | 49.512 | 11.653 | 1.00 10.00 | A_13 |
| MOTA | 300 | CE1 | PHE | 35 | 76.494 | 47.485 | 13.212 | 1.00 12.45 | A_13 |
| MOTA | 301 | CE2 | PHE | 35 | 76.224 | 49.538 | 12.002 | 1.00 17.92 | |
| ATOM | 302 | CZ | PHE | 35 | 75.684 | | | | A_13 |
| ATOM | 303 | C | | | | 48.525 | 12.777 | 1.00 13.29 | A_13 |
| | | | PHE | 35 | 82.170 | 47.754 | 12.236 | 1.00 11.31 | A_13 |
| MOTA | 304 | 0 | PHE | 35 | 82.493 | 46.573 | 12.034 | 1.00 11.37 | A_13 |
| MOTA | 305 | N | LYS | 36 | 82.962 | 48.778 | 11.945 | 1.00 17.06 | A_13 |
| MOTA | 307 | CA | LYS | 36 | 84.293 | 48.573 | 11.400 | 1.00 17.41 | |
| ATOM | 308 | CB | LYS | 36 | 84.991 | 49.922 | | | A_13 |
| ATOM | 309 | CG | LYS | 36 | | | 11.208 | 1.00 11.20 | A_13 |
| | | | | | 86.282 | 49.792 | 10.439 | 1.00 28.84 | A_13 |
| MOTA | 310 | CD | LYS | 36 | 87.246 | 50.917 | 10.738 | 1.00 24.52 | A_13 |
| ATOM | 311 | CE | LYS | 36 | 88.542 | 50.703 | 9.978 | 1.00 12.87 | A_13 |
| ATOM | 312 | NZ | LYS | 36 | 88.264 | 50.536 | 8.514 | 1.00 23.69 | A_13 |
| MOTA | 316 | C | LYS | 36 | 85.122 | 47.685 | 12.345 | 1.00 16.09 | W_T2 |
| ATOM | 317 | Ö | LYS | 36 | 85.701 | | | | A_13 |
| MOTA | 318 | N | | | | 46.686 | 11.938 | 1.00 21.50 | A_13 |
| | | | LYS | 37 | 85.173 | 48.057 | 13.613 | 1.00 12.42 | A_13 |
| ATOM | 320 | CA | LYS | 37 | 85.926 | 47.303 | 14.591 | 1.00 12.36 | A_13 |
| MOTA | 321 | CB | LYS | 37 | 85.953 | 48.066 | 15.917 | 1.00 13.65 | A_13 |
| MOTA | 322 | CG | LYS | 37 | 86.744 | 47.374 | 17.028 | 1.00 13.38 | |
| ATOM | 323 | CD | LYS | 37 | 88.192 | 47.125 | | | A_13 |
| ATOM | 324 | CE | LYS | | | | 16.616 | 1.00 38.32 | A_13 |
| | | | | 37 | 88.750 | 45.825 | 17.205 | 1.00 34.46 | A_13 |
| MOTA | 325 | NZ | LYS | 37 | 88.234 | 44.576 | 16.557 | 1.00 12.49 | A_13 |
| ATOM | 329 | C | LYS | 37 | 85.372 | 45.887 | 14.786 | 1.00 17.04 | A_13 |
| MOTA | 330 | 0 | LYS | 37 | 86.131 | 44.958 | 15.053 | 1.00 18.14 | A_13 |
| ATOM | 331 | N | ALA | 38 | 84.061 | 45.711 | 14.649 | 1.00 24.47 | |
| ATOM | 333 | CA | ALA | 38 | 83.452 | 44.392 | | | A_13 |
| ATOM | 334 | CB | ALÀ | 38 | | | 14.822 | 1.00 11.03 | A_13 |
| MOTA | 335 | C | | | 81.941 | 44.504 | 14.890 | 1.00 14.71 | A_13 |
| | | | ALA | 38 | 83.900 | | 13.697 | 1.00 20.27 | A_13 |
| MOTA | 336 | 0 | ALA | 38 | 84.143 | 42.266 | 13.936 | 1.00 18.80 | A_13 |
| MOTA | 337 | · N | PHE | 39 | 84.021 | 43.971 | 12.477 | 1.00 22.58 | A_13 |
| MOTA | 339 | CA | PHE | 39 | 84.492 | 43.158 | 11.355 | 1.00 18.87 | |
| MOTA | 340 | CB | PHE | 39 | 84.350 | 43.899 | 10.027 | | A_13 |
| ATOM | 341 | CG | PHE | 39 | | | | 1.00 19.91 | A_13 |
| MOTA | 342 | CD1 | | | 82.993 | 43.783 | 9.414 | 1.00 10.00 | A_13 |
| | | | | 39 | 82.266 | 44.915 | 9.097 | 1.00 17.54 | A_13 |
| ATOM | 343 | | PHE | 39 | 82.438 | 42.533 | 9.143 | 1.00 15.92 | A_13 |
| ATOM | 344 | CEI | PHE | 39 | 81.008 | 44.808 | 8.520 | 1.00 20.75 | A_13 |
| ATOM | 345 | CE2 | PHE | 39 | 81.186 | 42.418 | 8.569 | 1.00 10.00 | A_13 |
| ATOM | 346 | CZ | PHE | 39 | 80.467 | 43.555 | | 1.00 10.00 | |
| MOTA | 347 | С | PHE | 39 | 85.955 | 42.827 | | | A_13 |
| MOTA | 348 | Ö | PHE | 39 | | | 11.589 | 1.00 16.52 | A_13 |
| ATOM | 349 | N | | | 86.382 | 41.689 | 11.387 | 1.00 19.70 | . A_13 |
| MOTA | | | LYS | 40 | 86.699 | 43.822 | 12.072 | 1.00 21.31 | A_13 |
| | 351 | CA | LYS | 40 | 88.117 | 43.673 | 12.369 | 1.00 20.07 | A_13 |
| ATOM | 352 | CB | LYS | 40 | 88.703 | 44.967 | 12.927 | 1.00 13.77 | A_13 |
| MOTA | 353 | CG | LYS | 40 | 90.192 | 44.885 | 13.171 | 1.00 11.54 | A_13 |
| MOTA | 354 | CD | LYS | 40 | 90.757 | 46.242 | 13.507 | 1.00 10.34 | |
| MOTA | 355 | CE | LYS | 40 | 92.236 | 46.142 | | | A_13 |
| ATOM | 356 | NZ | LYS | 40 | 92.468 | | 13.838 | 1.00 11.24 | A_13 |
| ATOM | 360 | C | LYS | | | 45.518 | 15.179 | 1.00 27.33 | A_13 |
| ATOM | 361 | | | 40 | 88.352 | 42.534 | 13.337 | 1.00 12.06 | A_13 |
| | | 0 | LYS | 40 | 89.252 | 41.719 | 13.124 | 1.00 25.09 | A_13 |
| MOTA | 362 | N | VAL | 41 | 87.495 | 42.418 | 14.349 | 1.00 12.26 | A_13 |
| MOTA | 364 | CA | VAL | 41 | 87.630 | 41.331 | 15.325 | 1.00 17.89 | A_13 |
| MOTA | 365 | CB | VAL | 41 | 86.351 | 41.205 | 16.216 | 1.00 10.00 | |
| ATOM | 366 | CG1 | VAL | 41 | 86.298 | 39.865 | | | A_13 |
| MOTA | 367 | | VAL | 41 | | | 16.894 | 1.00 23.82 | A_13 |
| MOTA | 368 | C | | | 86.329 | 42.274 | 17.259 | 1.00 17.65 | A_13 |
| | | | VAL | 41 | 87.822 | 40.009 | 14.560 | 1.00 23.06 | A_13 |
| ATOM | 369 | 0 | VAL | 41 | 88.664 | 39.168 | 14.912 | 1.00 11.82 | A_13 |
| MOTA | 370 | N | TRP | 42 | 87.069 | 39.871 | 13.471 | 1.00 21.42 | A_13 |
| MOTA | 372 | CA | TRP | 42 | 87.085 | 38.666 | 12.661 | 1.00 21.32 | A_13 |
| ATOM | 373 | CB | TRP | 42 | 85.713 | 38:476 | 12.009 | 1.00 18.84 | |
| ATOM | 374 | CG | TRP | 42 | 84.605 | 38.387 | | | A_13 |
| ATOM | 375 | CD2 | | | | | 13.025 | 1.00 25.92 | A_13 |
| ATOM | 376 | | | 42 | 84.437 | 37.369 | 14.024 | 1.00 16.65 | A_13 |
| | | CE2 | | 42 | 83.260 | 37.680 | 14.737 | 1.00 17.58 | A_13 |
| MOTA | 377 | CE3 | | 42 | 85.165 | 36.223 | 14.380 | 1.00 11.14 | A_13 |
| MOTA | 378 | CD1 | | 42 | 83.563 | 39.249 | 13.179 | 1.00 10.00 | A_13 |
| MOTA | 379 | NEl | TRP | 42 | 82.755 | 38.832 | 14.200 | 1.00 10.91 | A_13 |
| MOTA | 381 | CZ2 | TRP | 42 | 82.785 | 36:879 | 15.793 | 1.00 10.91 | |
| MOTA | 382 | CZ3 | | 42 | 84.691 | 35.425 | | | A_13 |
| ATOM | 383 | CH2 | | 42 | 83.513 | | 15.436 | 1.00 23.68 | A_13 |
| ATOM | 384 | C | TRP | | | 35.759 | 16.125 | 1.00 12.75 | A_13 |
| | -04 | _ | INP | 42 | 88.190 | 38.600 | 11.623 | 1.00 27.45 | A_13 |

| ATOM 398 CR SER 43 88.413 39.702 19.891 1.00 29.46 A_13 ATOM 399 CR SER 43 89.422 39.769 9.891 1.00 19.65 A_13 ATOM 390 CR SER 43 89.452 39.769 9.891 1.00 19.65 A_13 ATOM 391 CR SER 43 91.758 39.119 9.834 1.00 17.99 A_13 ATOM 393 CR SER 43 91.758 39.119 9.834 1.00 17.99 A_13 ATOM 396 CR ASP 44 92.206 39.908 12.505 1.00 17.99 A_13 ATOM 397 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 398 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 398 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 399 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 399 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 399 CR ASP 44 92.206 39.908 12.505 1.00 16.90 A_13 ATOM 400 CR ASP 44 92.506 42.003 11.00 15.93 A_13 ATOM 401 CR ASP 44 92.506 42.003 11.00 15.93 A_13 ATOM 402 CR ASP 44 92.781 38.523 12.729 1.00 12.12 A_13 ATOM 403 N VAL 45 91.911 37.523 12.745 1.00 19.51 A_13 ATOM 405 CR VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 406 CR VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 408 CR VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 409 CR VAL 45 91.833 35.678 1.381 1.00 16.30 A_13 ATOM 409 CR VAL 45 91.833 35.678 1.381 1.00 16.30 A_13 ATOM 409 CR VAL 45 91.833 35.678 1.381 1.00 16.30 A_13 ATOM 409 CR VAL 45 91.833 35.678 1.381 1.00 16.30 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 24.13 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 24.13 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 24.13 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 24.13 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 12.433 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 24.13 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 12.433 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 12.433 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 12.433 A_13 ATOM 410 CR ASP 44 92.353 36.167 11.911 1.00 12.433 A_13 ATOM 410 CR ASP 44 92.354 36.454 | ATOM | 385 | 0 | TRP | 42 | 88.834 | 37.556 | 11.472 | 1.00 11.84 | A_13 |
|--|--------------|------------|---------|-----|----------|------------------|--------|--------|------------|--------------|
| ATOM 389 CB SER 43 89.342 00.993 8.991 1.00 16.16 4 7.13 ATOM 392 C SER 43 89.455 42.199 9.709 1.00 26.14 7.13 ATOM 392 C SER 43 90.837 39.615 10.691 1.00 11.53 7.13 ATOM 393 C SER 43 90.837 39.615 10.691 1.00 11.53 7.13 ATOM 394 CA A SEP 44 90.493 39.973 11.771 1.00 10.00 7.13 ATOM 395 CA SER 44 92.206 39.998 12.505 1.00 16.90 7.13 ATOM 397 CB ASP 44 92.206 39.998 12.505 1.00 16.90 7.13 ATOM 398 CA SEP 44 92.206 39.998 12.505 1.00 16.90 7.13 ATOM 399 CD SER 44 92.206 39.998 12.505 1.00 16.79 7.13 ATOM 399 CD SER 44 92.206 39.998 12.505 1.00 16.79 7.13 ATOM 399 CD SER 44 92.605 42.618 18.399 1.00 16.79 7.13 ATOM 400 CD SEP 44 92.814 42.533 12.754 1.00 19.50 7.13 ATOM 401 C SEP 44 92.874 42.533 12.754 1.00 19.50 7.13 ATOM 402 C ASP 44 92.874 42.533 12.754 1.00 19.50 7.13 ATOM 403 N VAL 45 91.31 37.523 12.745 1.00 20.21.21 ATOM 405 CA VAL 45 92.353 36.161 12.996 1.00 27.53 7.13 ATOM 407 CO1 VAL 45 92.353 36.161 12.996 1.00 27.53 7.13 ATOM 408 C CE VAL 45 91.83 35.678 14.381 1.00 16.30 7.13 ATOM 409 CO VAL 45 91.83 35.678 14.381 1.00 16.30 7.13 ATOM 401 C VAL 45 91.83 35.678 14.381 1.00 16.30 7.13 ATOM 401 C C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 403 C CA VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.828 35.877 14.931 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 45 91.848 33.578 12.157 1.00 10.00 8.13 ATOM 410 C VAL 46 91.898 38.30 9.00 9.00 9.00 9.00 9.00 9.00 9.00 9 | ATOM ATOM | 386 388 | N CA | SER | 43 43 | 88.413 89.449 | 39.702 | 10.909 | 1.00 25.46 | A_13 A_13 |
| ATOM 392 C SER 43 90.837 39.615 10.491 1.00 11.53 | | | | | | | | | | |
| ATOM 393 O SER 43 91.758 39.119 9.834 1.00 17.99 | | | | | | | | | | |
| ATOM 394 N ASP 44 90.949 39.973 11.771 1.00 10.00 A_13 ATOM 397 CB ASP 44 92.206 39.908 12.505 1.00 10.00 A_13 ATOM 398 CG ASP 44 92.206 39.908 12.505 1.00 17.79 A_13 ATOM 399 CG ASP 44 92.505 42.618 13.857 1.00 17.79 A_13 ATOM 399 CD ASP 44 92.505 42.618 14.920 1.00 17.21 A_13 ATOM 399 DO DASP 44 92.605 42.618 14.920 1.00 17.21 A_13 ATOM 401 C ASP 44 92.874 42.533 12.739 1.00 25.12 A_13 ATOM 401 C ASP 44 92.875 31.85.233 12.739 1.00 25.12 A_13 ATOM 402 C ASP 44 92.875 31.85.233 12.739 1.00 25.12 A_13 ATOM 405 CA ALL 45 93.956 38.523 12.739 1.00 26.12 A_13 ATOM 405 CA ALL 45 93.2133 31.523 12.739 1.00 26.12 A_13 ATOM 405 CB VALL 45 93.2133 31.523 12.739 1.00 27.83 A_13 ATOM 407 CG1 VAL 45 93.233 31.573 12.7495 1.00 17.83 A_13 ATOM 408 CG2 VAL 45 93.83 5.876 14.495 1.00 10.00 A_13 ATOM 409 C VAL 45 99.1.853 36.872 14.95 1.00 27.83 A_13 ATOM 401 O VAL 45 99.1.833 37.81 21.757 1.00 18.84 A_13 ATOM 410 C VAL 45 99.1.893 39.781 21.757 1.00 18.84 A_13 ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 411 N THR 46 89.750 35.705 10.694 1.00 16.30 A_13 ATOM 412 CB THR 46 89.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CG THR 46 89.750 35.705 10.694 1.00 16.30 A_13 ATOM 414 CB THR 46 89.750 35.705 10.694 1.00 16.30 A_13 ATOM 415 CG1 THR 46 89.7150 35.705 10.694 1.00 16.30 A_13 ATOM 412 CD PRO 47 91.693 31.805 7.705 10.00 10.00 1.53 A_13 ATOM 412 CD PRO 47 91.693 31.805 7.705 10.00 10.00 A_1.33 ATOM 412 CD PRO 47 91.693 31.805 7.705 10.00 10.00 A_1.33 ATOM 420 N PRO 47 92.099 35.416 5.805 1.00 10.00 10.99 A_1.34 ATOM 420 N PRO 47 92.099 35.416 5.805 1.00 10.00 1.00 A_1.34 ATOM 421 CD PRO 47 91.693 31.805 7.705 10.00 10.00 A_1.34 ATOM 422 CD PRO 47 91.693 31.805 7.714 1.00 11.00 A_1.34 ATOM 423 CD PRO 47 91.693 31.805 7.714 1.00 11.00 A_1.34 ATOM 424 CG PRO 47 92.099 35.416 5.805 1.00 21.40 A_1.34 ATOM 425 C PRO 47 92.099 35.416 5.805 1.00 21.40 A_1.34 ATOM 426 CD PRO 47 91.693 31.805 7.905 1.00 21.40 A_1.34 ATOM 427 N EEU 48 88.826 37.434 5.591 1.00 21.45 A_1.34 ATOM 426 CD PRO 47 91.695 31.806 6.808 1.10 0.10 0.0 | | | | | | | | | | |
| ATOM 396 CA ASP 44 92.066 39.908 12.505 1.00 16.90 | | | | | | | | | | |
| ATOM 398 CG ASP 44 92.544 42.013 13.839 1.00 15.93 | | 396 | CA | ASP | 44 | 92.206 | 39.908 | 12.505 | 1.00 16.90 | |
| ATOM 400 0D2 ASP 44 92.605 42.618 14.920 1.00 17.21 | | | | | | | | | | _ |
| ATOM 400 ODZ ASP 44 92.874 42.533 12.754 1.00 19.50 A_13 ATOM 401 C ASP 44 93.996 38.362 12.897 1.00 21.21 A_13 ATOM 403 N VAL 45 93.996 38.362 12.897 1.00 21.21 A_13 ATOM 405 CA VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 406 CB VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 407 CG1 VAL 45 92.853 15.678 14.381 1.00 10.00 A_13 ATOM 408 CG2 VAL 45 99.853 35.678 14.381 1.00 10.00 A_13 ATOM 409 C CVAL 45 99.853 35.877 14.955 1.00 22.03 A_13 ATOM 409 C VAL 45 99.853 35.877 14.955 1.00 10.866 A_13 ATOM 410 C VAL 45 99.863 33.978 12.157 1.00 18.84 A_13 ATOM 411 N THR 46 91.750 35.705 10.594 1.00 16.30 A_13 ATOM 411 N THR 46 91.750 35.705 10.594 1.00 16.30 A_13 ATOM 412 CB THR 46 89.128 36.004 9.682 1.00 13.14 48 A_13 ATOM 413 CG THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 414 CB THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 419 C THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 419 C THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 419 C THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 419 C THR 46 89.128 36.004 9.682 1.00 13.15 A_13 ATOM 412 CD PRO 47 91.683 35.416 5.815 1.00 17.51 A_13 ATOM 420 N PRO 47 91.683 35.416 5.815 1.00 17.51 A_13 ATOM 421 CD PRO 47 91.683 33.986 6.989 1.00 17.51 A_13 ATOM 422 CD PRO 47 92.099 35.416 5.815 1.00 17.57 A_13 ATOM 423 CB PRO 47 92.099 35.416 5.815 1.00 27.45 A_13 ATOM 424 CC PRO 47 91.095 36.788 4.116 1.00 17.57 A_13 ATOM 425 C PRO 47 91.095 36.788 4.116 1.00 17.57 A_13 ATOM 426 O PRO 47 91.095 36.788 4.116 1.00 17.57 A_13 ATOM 427 N LEU 48 88.826 37.434 5.581 1.00 27.45 A_13 ATOM 427 N LEU 48 88.826 37.434 5.581 1.00 22.09 A_13 ATOM 428 CD PRO 47 91.095 36.788 4.116 1.00 17.00 A_13.8 ATOM 429 CA LEU 48 88.826 37.434 5.581 1.00 22.09 A_13 ATOM 420 CB LEU 48 88.826 37.434 5.591 1.00 22.09 A_13 ATOM 421 CD PRO 47 91.095 36.788 4.116 1.00 17.00 A_13.8 ATOM 423 CD PRO 47 91.095 36.788 4.116 1.00 17.00 A_13.8 ATOM 426 C PRO 47 91.095 36.788 4.116 1.00 17.00 A_13.8 ATOM 427 N LEU 48 88.99.918 6.10 31.80 1.00 11.00 A_13.8 ATOM 428 C PRO 47 91.095 36.788 4.116 1.00 11.00 A_13 | | | | | | | | | | |
| ATOM 401 C ASP 44 92.781 38.523 12.729 1.00 26.12 A_13 ATOM 402 O ASP 44 93.996 38.362 12.897 1.00 21.21 A_13 ATOM 405 CA VAL 45 91.911 37.523 12.745 1.00 20.89 A_13 ATOM 406 CB VAL 45 91.933 36.161 12.996 1.00 27.53 A_13 ATOM 407 CG1 VAL 45 91.833 35.676 14.391 1.00 16.30 A_13 ATOM 408 CG2 VAL 45 91.833 35.676 14.391 1.00 10.00 A_13 ATOM 409 C VAL 45 91.833 35.676 14.391 1.00 10.86 A_13 ATOM 409 C VAL 45 91.983 35.877 11.495 1.00 10.86 A_13 ATOM 410 O VAL 45 91.864 33.978 12.157 1.00 18.84 A_13 ATOM 411 N THR 46 91.780 35.705 10.694 1.00 16.30 A_13 ATOM 412 CB THR 46 89.129 33.609 9.028 1.00 31.53 A_13 ATOM 413 CA THR 46 89.129 33.609 9.028 1.00 31.53 A_13 ATOM 415 OG1 THR 46 89.129 33.609 9.028 1.00 31.53 A_13 ATOM 415 OG1 THR 46 89.112 35.675 8.257 1.00 10.99 A_13 ATOM 415 OG1 THR 46 89.112 35.675 8.257 1.00 10.99 A_13 ATOM 419 O THR 46 91.716 35.575 8.257 1.00 10.19 A_13 ATOM 419 O THR 46 91.716 35.575 8.257 1.00 10.19 A_13 ATOM 420 N FRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 421 CB TRR 47 92.022 35.6764 8.255 1.00 17.64 A_13 ATOM 421 CB TRR 47 92.023 33.398 6.985 1.00 17.54 A_13 ATOM 422 CD FRO 47 92.069 35.416 5.815 1.00 17.54 A_13 ATOM 423 CB FRO 47 92.059 35.416 5.815 1.00 17.54 A_13 ATOM 420 N FRO 47 92.059 35.416 5.815 1.00 17.54 A_13 ATOM 421 CD FRO 47 99.958 36.786 8.55 1.00 17.54 A_13 ATOM 422 CB FRO 47 99.958 36.786 8.55 1.00 17.54 A_13 ATOM 423 CB FRO 47 99.958 36.786 8.55 1.00 17.54 A_13 ATOM 424 CB FRO 47 99.958 36.786 8.551 1.00 17.54 A_13 ATOM 425 CB FRO 47 99.958 36.786 8.551 1.00 17.54 A_13 ATOM 426 CB FRO 47 99.958 36.786 8.551 1.00 17.54 A_13 ATOM 427 N LEU 48 88.958 39.16 5.481 1.00 17.55 A_13 ATOM 433 CD LEU 48 88.958 39.16 5.481 1.00 17.55 A_13 ATOM 434 CB LEU 48 86.848 8.569 39.60 1.00 10.00 A_13 ATOM 435 CB FRO 47 99.958 36.786 4.116 1.00 11.08 A_13 ATOM 436 N ASN 49 89.968 41.159 3.885 1.00 12.14 A_13 ATOM 437 CD LEU 48 88.959 39.60 1.00 1.00 1.00 A_13 ATOM 436 CB PHE 50 88.868 41.705 7.7490 1.00 10.15.34 A_13 ATOM 446 CB PHE 50 88.868 41.705 7.7490 1.00 10.00 A_13 ATOM 457 CB | | | | | | | | | | |
| ATOM 403 N° VAL 45 91.911 37.523 12.745 1.00 20.89 Ala ATOM 406 CA VAL 45 92.353 36.161 12.996 1.00 27.53 Ala ATOM 407 CGI VAL 45 91.853 35.676 14.381 1.00 16.30 Ala ATOM 408 CCZ VAL 45 90.348 35.857 14.495 1.00 10.86 Ala ATOM 409 C VAL 45 91.983 35.867 14.495 1.00 10.86 Ala ATOM 409 C VAL 45 91.986 35.187 11.911 1.00 24.33 Ala ATOM 410 O VAL 45 91.864 33.978 12.157 1.00 18.84 Ala ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 Ala ATOM 413 CA THR 46 91.233 34.993 9.574 1.00 14.48 Ala ATOM 414 CB THR 46 89.279 33.609 9.028 1.00 31.53 Ala ATOM 415 OGI THR 46 89.279 33.609 9.028 1.00 31.53 Ala ATOM 415 OGI THR 46 89.175 36.014 9.00 10.99 Ala ATOM 418 C THR 46 89.112 35.755 8.257 1.00 25.10 Ala ATOM 419 O THR 46 91.716 35.755 8.257 1.00 15.34 Ala ATOM 419 O THR 46 92.022 36.764 8.256 1.00 17.64 Ala ATOM 420 N FRO 47 91.688 34.845 7.114 1.00 15.31 Ala ATOM 421 CD PRO 47 91.459 33.398 6.985 1.00 17.94 Ala ATOM 422 CA PRO 47 92.059 35.416 5.815 1.00 21.50 Ala ATOM 422 CB FRO 47 92.059 35.416 5.815 1.00 21.50 Ala ATOM 423 CB FRO 47 92.393 33.688 4.116 1.00 17.77 Ala ATOM 424 CG FRO 47 92.399 33.6182 4.911 1.00 17.57 Ala ATOM 425 CD PRO 47 90.991 36.248 5.256 1.00 11.94 Ala ATOM 427 N LEU 48 88.826 37.434 5.581 1.00 27.45 Ala ATOM 429 CA LEU 48 88.826 37.434 5.581 1.00 27.45 Ala ATOM 420 CD PRO 47 90.991 36.248 5.256 1.00 11.98 Ala ATOM 420 CD PRO 47 90.991 36.248 5.256 1.00 11.94 Ala ATOM 420 CD PRO 47 90.991 36.248 5.256 1.00 11.94 Ala ATOM 420 CD PRO 47 90.991 36.248 5.556 1.00 27.45 Ala ATOM 421 CD PRO 47 90.991 36.248 5.556 1.00 27.45 Ala ATOM 422 CB PRO 47 90.991 36.248 5.556 1.00 27.45 Ala ATOM 423 CB PRO 47 90.991 36.248 5.556 1.00 27.45 Ala ATOM 424 CG PRO 47 90.991 36.248 5.556 1.00 27.45 Ala ATOM 425 CD PRO 47 90.991 36.248 5.557 1.00 27.45 Ala ATOM 426 CD PRO 47 90.991 36.248 5.557 1.00 27.45 Ala ATOM 427 N LEU 48 88.991 36.576 6.432 1.00 10.00 Ala ATOM 428 CD PRO 47 90.991 36.248 5.557 1.00 27.45 Ala ATOM 429 CA LEU 48 86.891 88.991 80.557 6.018 1.00 10.00 Ala ATOM 430 CB LEU 48 86.891 88.991 80.00 10.00 | | | | | | | | | | A_13 |
| ATOM 405 CB VAL 45 92.353 36.161 12.996 1.00 27.53 A_13 ATOM 407 CGI VAL 45 92.857 35.678 14.381 1.00 16.30 A_13 ATOM 408 CG2 VAL 45 92.857 36.472 15.504 1.00 10.06 A_13 ATOM 409 CC VAL 45 91.928 35.877 11.911 1.00 24.33 A_13 ATOM 410 C VAL 45 91.928 35.877 11.911 1.00 24.33 A_13 ATOM 410 C VAL 45 91.928 35.877 11.911 1.00 16.30 A_13 ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 89.750 34.893 9.574 1.00 14.48 A_13 ATOM 414 CB THR 46 89.750 34.996 9.662 1.00 22.05 A_13 ATOM 415 CGI THR 46 89.129 36.014 90.040 1.00 10.99 A_13 ATOM 417 CGI THR 46 89.112 36.014 90.040 1.00 10.99 A_13 ATOM 419 O THR 46 91.716 35.575 8.257 1.00 25.10 A_13 ATOM 420 N PRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 421 CD PRO 47 91.689 33.398 6.985 1.00 17.94 A_13 ATOM 422 CA PRO 47 92.059 35.416 5.815 1.00 17.64 A_13 ATOM 422 CA PRO 47 92.059 33.938 6.985 1.00 17.94 A_13 ATOM 423 CB PRO 47 92.199 34.182 4.911 1.00 17.57 A_13 ATOM 424 CG PRO 47 92.199 36.748 5.848 1.00 27.45 A_13 ATOM 425 C PRO 47 99.199 36.6788 4.116 1.00 11.08 1.03 ATOM 426 C PRO 47 91.095 36.788 4.116 1.00 11.08 A_13 ATOM 427 N LEU 48 89.918 36.587 6.018 1.00 10.09 A_13 ATOM 428 CD LEU 48 88.826 37.434 5.858 1.00 10.00 A_13 ATOM 429 CA LEU 48 88.826 37.434 5.858 1.00 10.00 A_13 ATOM 430 CB LEU 48 88.826 39.366 6.085 1.00 10.00 A_13 ATOM 431 CG LEU 48 88.836 38.867 6.358 1.00 10.00 A_13 ATOM 432 CDI LEU 48 88.836 39.366 6.085 1.00 11.08 A_13 ATOM 432 CDI LEU 48 89.938 36.567 6.018 1.00 10.00 A_13 ATOM 433 CDI LEU 48 89.938 36.567 6.018 1.00 10.00 A_13 ATOM 434 C LEU 48 89.938 36.567 6.035 1.00 11.08 A_13 ATOM 435 C LEU 48 89.936 41.759 38.936 6.030 1.00 10.358 A_13 ATOM 436 N ASN 49 89.936 41.759 38.936 6.030 1.00 12.09 A_13 ATOM 437 C LEU 48 89.936 41.759 38.936 6.030 1.00 12.38 A_13 ATOM 436 N ASN 49 89.936 41.759 38.937 1.00 10.00 A_13 ATOM 437 C A BREE 50 88.838 49.936 6.030 1.00 10.00 A_13 ATOM 446 C A THR 51 86.939 89.936 41.216 6.395 1.00 10.22.56 A_13 ATOM 445 C B PHE 50 88.83 | | | - | | | | | | | |
| ATOM 406 CB VAL 45 91.833 35.678 14.381 1.00 16.30 | | | | | | | | | | |
| ATOM 409 C VAL 45 90.348 35.857 14.495 1.00 10.86 A_13 ATOM 410 O VAL 45 91.964 33.978 12.157 1.00 18.30 A_13 ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 89.1750 33.4796 9.662 1.00 22.05 A_13 ATOM 415 OG1 THR 46 89.1750 34.796 9.662 1.00 22.05 A_13 ATOM 415 OG1 THR 46 89.1716 35.575 8.257 1.00 25.10 A_13 ATOM 417 CG2 THR 46 89.1716 35.575 8.257 1.00 25.10 A_13 ATOM 419 O THR 46 92.022 36.764 8.256 1.00 11.53 A_13 ATOM 419 O THR 46 92.022 36.764 8.256 1.00 17.54 A_13 ATOM 420 N PRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 421 CD PRO 47 91.688 33.398 6.985 1.00 17.54 A_13 ATOM 422 CA PRO 47 92.199 34.162 4.911 1.00 17.57 A_13 ATOM 423 CB PRO 47 92.199 34.162 4.911 1.00 17.57 A_13 ATOM 424 CG PRO 47 92.299 34.162 5.815 1.00 21.50 A_13 ATOM 425 C PRO 47 92.369 33.041 5.848 1.00 27.45 A_13 ATOM 426 O PRO 47 90.991 36.348 5.256 1.00 21.10 A_13 ATOM 427 N LEU 48 89.918 36.567 6.018 1.00 11.08 A_13 ATOM 429 CA LEU 48 88.926 37.434 5.581 1.00 11.08 A_13 ATOM 429 CA LEU 48 88.926 37.434 5.581 1.00 12.09 A_13 ATOM 431 CG LEU 48 88.938 36.567 6.018 1.00 11.08 A_13 ATOM 432 CD LEU 48 88.938 36.567 6.018 1.00 11.08 A_13 ATOM 432 CD LEU 48 88.938 36.567 6.018 1.00 12.00 A_13 ATOM 431 CG LEU 48 88.938 36.567 6.018 1.00 12.00 A_13 ATOM 432 CD LEU 48 88.938 37.38 41.112 4.00 15.92 A_13 ATOM 430 CB LEU 48 89.918 36.567 6.018 1.00 12.00 A_13 ATOM 431 CG LEU 48 88.938 36.567 6.018 1.00 12.00 A_13 ATOM 432 CD LEU 48 88.938 37.38 41.112 4.00 15.93 A_13 ATOM 430 CB LEU 48 89.918 36.567 6.018 1.00 12.00 A_13 ATOM 431 CG LEU 48 86.848 35.867 6.038 1.00 12.00 A_13 ATOM 432 CD LEU 48 86.848 35.867 6.038 1.00 12.00 A_13 ATOM 430 CB LEU 48 89.918 39.666 5.167 1.00 16.47 A_13 ATOM 431 CG LEU 48 86.848 37.866 5.167 1.00 16.47 A_13 ATOM 432 CD LEU 48 86.848 37.866 5.666 5.167 1.00 16.647 A_13 ATOM 430 CB LEU 48 86.848 37.866 6.628 1.00 10.00 A_13 ATOM 430 CB LEU 48 89.918 6.00 A_13 ATOM 430 CD ARS 49 90.90 A_13 ATOM 430 CD ARS 49 90.90 A_13 ATOM 430 CD ARS | | | | | | | | | | A_13 |
| ATOM 409 C VAL 45 91.928 35.187 11.911 1.00 24.33 A_13 ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 91.293 34.893 9.574 1.00 14.48 A_13 ATOM 414 CB THR 46 89.279 33.609 9.562 1.00 22.05 A_13 ATOM 415 061 THR 46 89.279 33.609 9.562 1.00 22.05 A_13 ATOM 417 C62 THR 46 89.279 33.609 9.028 1.00 10.09 A_13 ATOM 418 C THR 46 89.279 33.609 9.028 1.00 10.09 A_13 ATOM 419 0 THR 46 91.716 35.575 8.257 1.00 25.10 1.13 ATOM 419 C THR 46 91.716 35.575 8.257 1.00 25.10 1.13 ATOM 420 N PRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 421 CD PRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 422 CA PRO 47 92.069 35.416 5.815 1.00 17.94 A_13 ATOM 423 CB PRO 47 92.069 35.416 5.815 1.00 17.94 A_13 ATOM 424 CG PRO 47 92.199 34.182 4.911 1.00 17.57 A_13 ATOM 425 C PRO 47 90.991 35.041 5.848 1.00 27.45 A_13 ATOM 426 O PRO 47 90.991 36.348 5.256 1.00 17.57 A_13 ATOM 427 N LEU 48 89.918 36.557 6.018 1.00 11.08 A_13 ATOM 429 CA LEU 48 88.826 37.434 5.581 1.00 12.00 A_13 ATOM 430 CB LEU 48 88.826 37.434 5.581 1.00 12.00 A_13 ATOM 431 CG LEU 48 88.918 35.557 6.018 1.00 11.08 A_13 ATOM 432 CD LEU 48 88.918 35.557 6.018 1.00 11.08 A_13 ATOM 433 CD LEU 48 88.918 35.857 6.018 1.00 11.08 A_13 ATOM 434 C LEU 48 88.918 35.857 6.018 1.00 11.08 A_13 ATOM 431 CG LEU 48 88.918 35.857 6.018 1.00 11.08 A_13 ATOM 432 CD LEU 48 88.936 39.670 6.482 1.00 15.58 A_13 ATOM 433 CD LEU 48 88.936 39.670 6.482 1.00 15.58 A_13 ATOM 434 C LEU 48 89.918 36.557 6.018 1.00 12.00 A_13 ATOM 435 O LEU 48 89.918 36.957 6.432 1.00 15.92 A_13 ATOM 436 CA ASN 49 89.936 39.666 6.480 1.00 17.28 A_13 ATOM 437 C LEU 48 89.918 39.967 6.432 1.00 12.02 2.55 A_13 ATOM 436 CA ASN 49 89.936 39.670 6.492 1.00 15.02 2.55 A_13 ATOM 436 CA ASN 49 89.936 41.569 1.00 12.00 2.77 A_13 ATOM 446 O ASN 49 89.936 41.569 1.00 12.00 2.1.73 A_13 ATOM 446 CA PHE 50 85.738 41.112 4.717 1.00 26.84 A_13 ATOM 446 CA PHE 50 85.738 41.128 3.400 1.00 21.47 A_13 ATOM 446 CA PHE 50 85.866 41.705 1.00 1.00 1.00 2.1.13 ATOM 465 CC PHE 50 85.866 41.705 1.00 1.00 1.00 2.1.13 ATOM 465 CC | | | | | | | | | | |
| ATOM 410 0 VAL 45 91.864 33.978 12.157 1.00 18.84 A_13 ATOM 411 N THR 46 91.750 35.705 10.694 1.00 16.30 A_13 ATOM 413 CA THR 46 91.750 34.796 9.662 1.00 12.05 A_13 ATOM 414 CB THR 46 89.750 34.796 9.662 1.00 12.05 A_13 ATOM 415 CG1 THR 46 89.713 36.014 9.040 1.00 10.09 A_13 ATOM 417 CG2 THR 46 89.113 36.014 9.040 1.00 10.09 A_13 ATOM 419 CC THR 46 99.1716 35.575 8.257 1.00 25.10 A_13 ATOM 419 O THR 46 99.1716 35.575 8.257 1.00 25.10 A_13 ATOM 419 O THR 46 99.022 36.764 8.256 1.00 17.04 A_13 ATOM 420 N PRO 47 91.688 34.845 7.114 1.00 15.31 A_13 ATOM 421 CD PRO 47 91.489 33.398 6.985 1.00 17.50 A_13 ATOM 422 CA PRO 47 92.069 35.416 5.815 1.00 21.50 A_13 ATOM 424 CG PRO 47 92.069 35.416 5.815 1.00 21.50 A_13 ATOM 425 C PRO 47 92.069 35.416 5.815 1.00 21.50 A_13 ATOM 426 O PRO 47 92.069 33.041 5.848 1.00 27.45 A_13 ATOM 427 N LEU 48 89.918 36.567 6.018 1.00 17.97 A_13 ATOM 427 N LEU 48 88.826 37.434 5.581 1.00 22.04 A_13 ATOM 430 CB LEU 48 88.826 37.434 5.581 1.00 22.05 A_13 ATOM 431 CG LEU 48 86.033 35.678 6.018 1.00 10.00 A_13 ATOM 432 CD1 LEU 48 88.826 37.434 5.581 1.00 22.05 A_13 ATOM 433 CD2 LEU 48 86.033 35.676 6.018 1.00 15.92 A_13 ATOM 434 C LEU 48 86.931 35.676 6.435 1.00 22.590 A_13 ATOM 435 C LEU 48 86.931 35.616 5.157 1.00 16.59 A_13 ATOM 436 N ASN 49 88.569 39.670 4.723 1.00 15.92 A_13 ATOM 437 CD LEU 48 89.938 41.116 1.00 11.00 A_13 ATOM 438 CA ASN 49 88.755 37.212 6.432 1.00 15.92 A_13 ATOM 436 C ASN 49 89.936 41.569 3.885 1.00 12.20 A_13 ATOM 437 CD LEU 48 89.938 41.116 1.00 22.09 A_13 ATOM 436 C ASN 49 88.769 39.670 4.723 1.00 26.12 A_13 ATOM 437 C D BU EU 48 89.938 41.116 1.00 21.20 A_13 ATOM 436 C ASN 49 89.936 41.569 3.885 1.00 18.29 A_13 ATOM 437 C D BU EU 48 89.936 41.259 1.00 16.01 A_13 ATOM 446 C ASN 49 89.936 41.569 3.885 1.00 18.29 A_13 ATOM 447 N PHE 50 86.732 41.128 3.400 1.00 20.217 A_13 ATOM 446 C ASN 49 89.936 41.690 1.00 0.00 A_13 ATOM 447 N PHE 50 86.732 41.128 3.400 1.00 20.217 A_13 ATOM 446 C ASN 49 89.936 41.690 3.9967 0.00 1.00 21.13 A_13 ATOM 446 C ASN 49 89.936 41.569 3.895 1 | | | | | • | | | | | |
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| ATOM 414 CB THR 46 89.750 34.796 9.662 1.00 22.05 | | | - | | | | | | | |
| ATOM 417 CC2 THR 46 89.112 36.014 9.040 1.00 10.99 | | | | | | | | | | |
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| ATOM 436 N ASN 49 88.569 39.670 4.723 1.00 26.12 A_13 ATOM 438 CA ASN 49 88.738 41.112 4.717 1.00 26.84 A_13 ATOM 439 CB ASN 49 89.936 41.569 3.885 1.00 18.29 A_13 ATOM 440 CG ASN 49 90.010 40.912 2.568 1.00 22.55 A_13 ATOM 441 OD1 ASN 49 90.928 40.131 2.305 1.00 24.41 A_13 ATOM 442 ND2 ASN 49 89.068 41.235 1.693 1.00 46.51 A_13 ATOM 445 C ASN 49 87.416 41.705 4.259 1.00 12.18 A_13 ATOM 446 O ASN 49 86.732 41.128 3.400 1.00 20.77 A_13 ATOM 447 N PHE 50 87.025 42.802 4.900 1.00 21.39 A_13 ATOM 449 CA PHE 50 85.738 43.439 4.642 1.00 10.00 A_13 ATOM 450 CB PHE 50 84.914 43.440 5.932 1.00 11.45 A_13 ATOM 451 CG PHE 50 84.863 42.098 6.629 1.00 10.63 A_13 ATOM 452 CD1 PHE 50 85.886 41.705 7.490 1.00 10.00 A_13 ATOM 453 CD2 PHE 50 85.886 41.705 7.490 1.00 10.00 A_13 ATOM 455 CE2 PHE 50 83.809 41.216 6.395 1.00 14.63 A_13 ATOM 455 CE2 PHE 50 83.809 41.216 6.395 1.00 14.63 A_13 ATOM 455 CD2 PHE 50 85.886 40.457 8.097 1.00 26.88 A_13 ATOM 456 CZ PHE 50 84.801 39.581 7.852 1.00 10.30 A_13 ATOM 457 C PHE 50 85.867 44.842 4.093 1.00 22.56 A_13 ATOM 458 O PHE 50 86.638 45.644 4.612 1.00 19.33 A_13 ATOM 459 N THR 51 85.099 45.129 3.044 1.00 21.47 A_13 ATOM 465 CB THE 51 85.602 46.336 0.895 1.00 15.39 A_13 ATOM 465 CG THR 51 85.602 46.336 0.895 1.00 15.39 A_13 ATOM 465 CG THR 51 85.602 46.336 0.895 1.00 15.39 A_13 ATOM 465 CG THR 51 85.602 46.306 0.895 1.00 15.39 A_13 ATOM 465 CG THR 51 85.602 46.433 2.371 1.00 24.21 A_13 ATOM 465 CG THR 51 85.602 46.433 2.371 1.00 24.21 A_13 ATOM 466 C THR 51 85.602 46.306 0.895 1.00 15.39 A_13 ATOM 467 O THR 51 85.602 46.306 0.895 1.00 22.56 A_13 ATOM 468 N ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 470 CA ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 471 CB ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 473 CD ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | | | | | | | | | |
| ATOM 438 CA ASN 49 88.738 41.112 4.717 1.00 26.84 A_13 ATOM 439 CB ASN 49 99.010 40.912 2.568 1.00 18.29 A_13 ATOM 441 OD1 ASN 49 90.010 40.912 2.568 1.00 22.55 A_13 ATOM 442 ND2 ASN 49 90.928 40.131 2.305 1.00 24.41 A_13 ATOM 442 ND2 ASN 49 89.068 41.235 1.693 1.00 46.51 A_13 ATOM 445 C ASN 49 87.416 41.705 4.259 1.00 12.18 A_13 ATOM 446 O ASN 49 86.732 41.128 3.400 1.00 20.77 A_13 ATOM 447 N PHE 50 87.025 42.802 4.900 1.00 21.39 A_13 ATOM 449 CA PHE 50 85.738 43.439 4.642 1.00 10.00 A_13 ATOM 450 CB PHE 50 84.863 42.098 6.629 1.00 11.45 A_13 ATOM 451 CG PHE 50 84.863 42.098 6.629 1.00 10.63 A_13 ATOM 453 CD2 PHE 50 85.886 41.705 7.490 1.00 10.00 A_13 ATOM 454 CE1 PHE 50 85.886 41.705 7.490 1.00 10.00 A_13 ATOM 455 CE2 PHE 50 83.809 41.216 6.395 1.00 14.63 A_13 ATOM 455 CE2 PHE 50 85.858 40.457 8.097 1.00 26.88 A_13 ATOM 455 CE2 PHE 50 85.866 41.705 7.852 1.00 10.30 A_13 ATOM 457 C PHE 50 85.867 44.842 4.093 1.00 22.56 A_13 ATOM 458 O PHE 50 86.638 45.644 4.612 1.00 19.33 A_13 ATOM 457 C PHE 50 85.867 44.842 4.093 1.00 22.56 A_13 ATOM 458 O PHE 51 85.099 45.129 3.044 1.00 21.47 A_13 ATOM 461 CA THR 51 85.099 45.129 3.044 1.00 21.47 A_13 ATOM 462 CB THR 51 85.602 46.306 0.895 1.00 15.39 A_13 ATOM 463 OG1 THR 51 85.602 46.306 0.895 1.00 24.21 A_13 ATOM 466 C THR 51 85.602 46.306 0.895 1.00 24.21 A_13 ATOM 466 C THR 51 85.602 46.306 0.895 1.00 22.56 A_13 ATOM 466 C THR 51 85.602 46.306 0.895 1.00 22.56 A_13 ATOM 466 C THR 51 85.602 46.306 0.895 1.00 25.47 A_13 ATOM 466 N ARG 52 82.393 49.004 2.871 1.00 20.53 A_13 ATOM 466 N ARG 52 82.393 49.004 2.871 1.00 20.53 A_13 ATOM 467 O THR 51 82.766 46.421 1.912 1.00 20.53 A_13 ATOM 468 N ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 470 CA ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 471 CB ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 473 CD ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 474 NE ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 474 NE ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 | | | | | | | | | | |
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| ATOM 467 O THR 51 82.766 46.421 1.912 1.00 20.53 A_13 ATOM 468 N ARG 52 83.653 48.294 2.797 1.00 16.53 A_13 ATOM 470 CA ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 471 CB ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 472 CG ARG 52 81.201 50.778 4.259 1.00 12.47 A_13 ATOM 473 CD ARG 52 81.462 51.879 5.278 1.00 19.61 A_13 ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | | | | | | | | | |
| ATOM 470 CA ARG 52 82.393 49.004 2.871 1.00 10.00 A_13 ATOM 471 CB ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 472 CG ARG 52 81.201 50.778 4.259 1.00 12.47 A_13 ATOM 473 CD ARG 52 81.462 51.879 5.278 1.00 19.61 A_13 ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | 467 | 0 | THR | 51 | 82.766 | 46.421 | 1.912 | 1.00 20.53 | A_13 |
| ATOM 471 CB ARG 52 82.490 50.085 3.939 1.00 10.00 A_13 ATOM 472 CG ARG 52 81.201 50.778 4.259 1.00 12.47 A_13 ATOM 473 CD ARG 52 81.462 51.879 5.278 1.00 19.61 A_13 ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | | | | | | | | | A_13 |
| ATOM 472 CG ARG 52 81.201 50.778 4.259 1.00 12.47 A_13 ATOM 473 CD ARG 52 81.462 51.879 5.278 1.00 19.61 A_13 ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | | | | | | | | | |
| ATOM 474 NE ARG 52 80.371 52.836 5.333 1.00 30.55 A_13 | | | CG | ARG | 52 | 81.201 | 50.778 | 4.259 | 1.00 12.47 | A_13 |
| | | | | | | | | | | |
| | | | | | | | | | | |

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| MOTA | 477 | NH1 A | ARG | 52 | 81.661 | 54.508 | 6.257 | 1.00 21.24 | מו א |
|--------|-----|--------------|-------|------|--------|--------|--------|------------|--------|
| ATOM | 480 | | | | | | | | A_13 |
| | | | ARG | 52 | 79.421 | 54.862 | 5.829 | 1.00 27.78 | A_13 |
| MOTA | 483 | | ARG | 52 | 81.980 | 49.620 | 1.540 | 1.00 30.22 | A_13 |
| ATOM | 484 | 0 7 | ARG | 52 | 82.782 | 50.269 | 0.859 | 1.00 16.27 | A_13 |
| MOTA | 485 | N I | LEU | 53 | 80.730 | 49.372 | 1.161 | 1.00 21.07 | A_13 |
| ATOM | 487 | | LEU | 53 | 80.159 | 49.914 | -0.062 | 1.00 15.73 | |
| | | | | | | | | | A_13 |
| ATOM | 488 | | LEU | 53 | 79.435 | 48.831 | -0.868 | 1.00 11.53 | A_13 |
| MOTA | 489 | CG I | LEU | 53 | 80.304 | 47.770 | -1.530 | 1.00 10.00 | A_13 |
| MOTA | 490 | CD1 | LEU | 53 | 79:429 | 46.790 | -2.296 | 1.00 13.21 | A_13 |
| MOTA | 491 | CD2 | | 53 | 81.280 | 48.443 | -2.448 | 1.00 12.78 | |
| | | | | | | | | | A_13 |
| MOTA | 492 | | LEU | 53 | 79.149 | 50.932 | 0.421 | 1.00 10.00 | A_13 |
| MOTA | 493 | | LEU | 53 | 78.463 | 50.713 | 1.411 | 1.00 13.62 | A_13 |
| MOTA | 494 | \mathbf{N} | HIS | 54 | 79.043 | 52.041 | -0.283 | 1.00 15.73 | A_13 |
| MOTA | 496 | CA I | HIS . | 54 | 78.102 | 53.065 | 0.126 | 1.00 12.47 | A_13 |
| MOTA | 497 | | HIS | 54 | 78.765 | 54.435 | 0.011 | 1.00 15.18 | |
| | 498 | | | | | | | | A_13 |
| ATOM | | | HIS | 54 | 79.967 | 54.589 | 0.884 | 1.00 21.27 | A_13 |
| MOTA | 499 | | HIS | 54 | 81.207 | 54.056 | 0.798 | 1.00 25.30 | A_13 |
| ATOM | 500 | ND1 | HIS | 54 | 79.951 | 55.338 | 2.043 | 1.00 16.48 | A_13 |
| MOTA | 502 | CE1 | HIS | 54 | 81.127 | 55.255 | 2.633 | 1.00 21.62 | A_13 |
| ATOM | 503 | | HIS | 54 | 81.910 | 54.482 | 1.899 | 1.00 29.91 | |
| ATOM | 505 | | | 54 | | | | | A_13 |
| | | | HIS | | 76.796 | 53.044 | -0.664 | 1.00 15.50 | A_13 |
| MOTA | 506 | • | HIS | 54 | 75.914 | 53.849 | -0.403 | 1.00 21.80 | A_13 |
| ATOM | 507 | N . | ASP | 55 | 76.707 | 52.178 | -1.671 | 1.00 18.31 | A_13 |
| MOTA | 509 | CA . | ASP | 55 | 75.509 | 52.077 | -2.502 | 1.00 17.23 | A_13 |
| MOTA | 510 | CB . | ASP | 55 | 75.645 | 52.928 | -3.773 | 1.00 19.94 | A_13 |
| MOTA | 511 | | ASP | 55 | 75.864 | 54.393 | -3.495 | | |
| ATOM | 512 | | | | | | | | A_13 |
| | | | ASP | 55 | 75.059 | 54.991 | -2.741 | 1.00 35.97 | A_13 |
| MOTA | 513 | OD2 | _ | 55 | 76.839 | 54.948 | -4.058 | 1.00 25.09 | A_13 |
| ATOM | 514 | C . | ASP | 55 | 75.343 | 50.645 | ~2.970 | 1.00 21.50 | A_13 |
| MOTA | 515 | 0 | ASP | 55 | 76.286 | 49.862 | -2.929 | 1.00 17.45 | A_13 |
| MOTA | 516 | | GLY | 56 | 74.160 | 50.337 | -3.489 | 1.00 10.31 | |
| ATOM | 518 | | GLY | 56 | | | | | A_13 |
| | | | | | 73.897 | 49.014 | -4.014 | 1.00 13.67 | A_13 |
| MOTA | 519 | | GLY | 56 | 73.842 | 47.869 | -3.030 | 1.00 17.61 | A_13 |
| MOTA | 520 | 0 | GLY | 56 | 73.683 | 48.065 | -1.825 | 1.00 12.57 | A_13 |
| ATOM | 521 | N | ILE | 57 | 73.943 | 46.653 | -3.560 | 1.00 22.27 | A_13 |
| ATOM . | 523 | CA | ILE | 57 | 73.895 | 45.460 | -2.737 | 1.00 11.39 | A_13 |
| MOTA | 524 | | ILE | 57 | 72.941 | 44.391 | -3.347 | 1.00 22.87 | |
| | 525 | | | | | | | | A_13 |
| MOTA | | | ILE | 57 | 73.365 | 42.995 | -2.955 | 1.00 22.98 | A_13 |
| MOTA | 526 | | ILE | . 57 | 71.522 | 44.582 | -2.787 | 1.00 30.87 | A_13 |
| MOTA | 527 | CD1 | ILE | 57 | 71.002 | 46.022 | -2.796 | 1.00 28.15 | A_13 |
| MOTA | 528 | C | ILE | 57 | 75.289 | 44.919 | -2.446 | 1.00 22.32 | A_13 |
| MOTA | 529 | 0 | ILE | 57 | 76.140 | 44.849 | -3.332 | 1.00 25.00 | A_13 |
| MOTA | 530 | | ALA | 58 | 75.517 | 44.631 | -1.168 | 1.00 25.02 | |
| ATOM | 532 | | ALA | | | | | _ | A_13 |
| | | | | 58 | 76.773 | 44.105 | -0.669 | 1.00 15.45 | A_13 |
| MOTA | 533 | | ALA | 58 | 77.366 | 45.060 | 0.358 | 1.00 11.62 | A_13 |
| MOTA | 534 | | ALA | 58 | 76.438 | 42.780 | -0.006 | 1.00 12.08 | A_13 |
| MOTA | 535 | 0 | ALA | 58 | 75.289 | 42.521 | 0.307 | 1.00 13.30 | A_13 |
| MOTA | 536 | N | ASP | 59 | 77.449 | 41.968 | 0.247 | 1.00 14.79 | A_13 |
| MOTA | 538 | CA | ASP | 59 | 77.245 | 40.675 | 0.880 | 1.00 18.50 | A_13 |
| MOTA | 539 | | ASP | 59 | 78.608 | 39.974 | 1.093 | 1.00 10.83 | |
| ATOM | | | | | | | | | A_13 |
| | 540 | | ASP | 59 | 79.425 | 39.858 | -0.210 | 1.00 23.35 | A_13 . |
| ATOM | 541 | | ASP | 59 | 80.598 | 40.266 | -0.236 | 1.00 17.98 | A_13 |
| MOTA | 542 | OD2 | ASP | 59 | 78.896 | 39.379 | -1.230 | 1.00 16.89 | A_13 |
| MOTA | 543 | C | ASP | 59 | 76.480 | 40.806 | 2.200 | 1.00 13.69 | A_13 |
| ATOM | 544 | 0 | ASP | 59 | 75.402 | 40.227 | 2.380 | 1.00 15.93 | A_13 |
| ATOM | 545 | N | ILE | 60 | 77.025 | | 3.109 | 1.00 13.15 | A_13 |
| ATOM | 547 | CA | ILE | 60 | 76.422 | 41.800 | 4.412 | 1.00 12.20 | |
| ATOM | 548 | CB | ILE | 60 | | | | | A_13 |
| | | _ | | | 77.500 | 41.695 | 5.508 | 1.00 12.12 | A_13 |
| ATOM | 549 | CG2 | | 60 | 76.921 | 42.060 | 6.864 | 1.00 19.27 | A_13 |
| MOTA | 550 | CGl | ILE | 60 | 78.118 | 40.287 | 5.481 | 1.00 10.00 | A_13 |
| ATOM | 551 | CD1 | ILE | 60 | 79.330 | 40.120 | 6.360 | 1.00 10.00 | A_13 |
| MOTA | 552 | С | ILE | 60 | 75.743 | | 4.456 | 1.00 17.78 | A_13 |
| ATOM | 553 | ŏ | ILE | 60 | 76.410 | | | | |
| | | | | | | | 4.478 | 1.00 18.65 | A_13 |
| ATOM | 554 | N | MET | 61 | 74.416 | | 4.431 | 1.00 12.54 | A_13 |
| ATOM | 556 | | MET | 61 | 73.640 | | 4.476 | 1.00 12.86 | A_13 |
| ATOM | 557 | CB | MET | 61 | 72.385 | | 3.604 | 1.00 18.16 | A_13 |
| MOTA | 558 | * CG | MET | 61 | 72.634 | 43.979 | 2.141 | 1.00 10.00 | A_13 |
| MOTA | 559 | SD | MET | 61 | 73.374 | | 1.251 | 1.00 10.69 | A_13 |
| ATOM | 560 | CE | MET | 61 | 71.836 | | 0.764 | | A_13 |
| ATOM | 561 | C | MET | 61 | 73.239 | | | | |
| ATOM | | | | | | | 5.921 | 1.00 10.15 | A_13 |
| | 562 | 0 | MET | 61 | 72.584 | | 6.547 | 1.00 18.13 | A_13 |
| ATOM | 563 | N | ILE | 62 | 73.706 | | 6.456 | 1.00 15.60 | A_13 |
| MOTA | 565 | CA | ILE | 62 | 73.452 | | 7.837 | 1.00 18.55 | A_13 |
| ATOM | 566 | CB | ILE | 62 | 74.723 | 46.828 | 8.437 | 1.00 10.00 | A_13 |
| MOTA | 567 | CG2 | ILE | 62 | 74.498 | | 9.900 | | A_13 |
| ATOM | 568 | | ILE | 62 | 75.936 | | | 1.00 11.04 | A_13 |
| ATOM | 569 | | ILE | 62 | 77.228 | | 8.891 | 1.00 10.00 | A_13 |
| | | | | | | | - · | | ~_±J |

| ATOM | 570 | С | ILE | 62 | 72 200 | 47 175 | 7 000 | 1 00 17 00 | |
|--------------|------------|-----------|------------|----------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 571 | | ILE | 62 62 | 72.289 72.335 | 47.172 | 7.920 | 1.00 17.99 | A_13 |
| ATOM | 572 | | SER | 63 | 71.285 | 48.208 46.896 | 7.264 8.751 | 1.00 12.72 1.00 10.00 | A_13 |
| ATOM | 574 | | SER | 63 | 70.149 | 47.803 | 8.882 | 1.00 10.00 | A_13 A_13 |
| ATOM | 575 | | SER | 63 | 69.016 | 47.364 | 7.956 | 1.00 13.06 | A_13 |
| ATOM | 576 | | SER | 63 | 68.448 | 46.146 | 8.415 | 1.00 27.90 | A_13 |
| MOTA | 57B | C | SER | 63 | 69.625 | 47.854 | 10.314 | 1.00 13.14 | A_13 |
| ATOM | 579 | Ö | SER | 63 | 69.869 | 46.951 | 11.101 | 1.00 22.10 | A_13 |
| ATOM | 580 | N | PHE | 64 | 68.919 | 48.932 | 10.640 | 1.00 21.17 | A_13 |
| ATOM | 582 | CA | PHE | 64 | 68.317 | 49.139 | 11.954 | 1.00 22.01 | A_13 |
| ATOM | 583 | CB | PHE | 64 | 68.777 | 50.468 | 12.574 | 1.00 10.98 | A_13 |
| ATOM | 584 | CG | PHE | 64 | 70.189 | 50.448 | 13.092 | 1.00 10.00 | A_13 |
| MOTA | 585 | CD1 | PHE | 64 | 70.473 | 49.885 | 14.322 | 1.00 10.00 | A_13 |
| MOTA | 586 | CD2 | PHE | 64 | 71.229 | 51.016 | 12.357 | 1.00 16.56 | A_13 |
| MOTA | 587 | CE1 | PHE | 64 | 71.777 | 49.885 | 14.825 | 1.00 10.00 | A_13 |
| MOTA | 588 | CE2 | PHE | 64 | 72.540 | 51.025 | 12.846 | 1.00 10.00 | A_13 |
| ATOM | 589 | CZ | PHE | 64 | 72.812 | 50.459 | 14.081 | 1.00 18.83 | A_13 |
| ATOM | 590 | C | PHE | 64 | 66.825 | 49.207 | 11.675 | 1.00 22.55 | A_13 |
| ATOM | 591 | 0 | PHE | 64 | 66.405 | 49.940 | 10.779 | 1.00 19.49 | A_13 |
| ATOM | 592 | N | GLY | 65 65 | 66.031 | 48.485 | 12.453 | 1.00 13.69 | A_13 |
| MOTA | 594 595 | CA | GLY | 65 65 | 64.593 | 48.491 | 12.238 | 1.00 10.70 | A_13 |
| MOTA MOTA | 595 596 | C O | GLY GLY | 65 65 | 63.894 64.559 | 48.138 | 13.521 | 1.00 12.62 | A_13 |
| ATOM | 597 | N | ILE | 66 | 62.577 | 47.777 48.309 | 14.491 13.565 | 1.00 18.29 1.00 13.69 | A_13 |
| ATOM | 599 | CA | ILE | 66 | 61.803 | 47.968 | 14.760 | 1.00 21.58 | A_13 A_13 |
| MOTA | 600 | CB | ILE | 66 | 61.227 | 49.228 | 15.503 | 1.00 30.51 | A_13 |
| MOTA | 601 | CG2 | ILE | 66 | 62.351 | 50.110 | 16.025 | 1.00 10.43 | A_13 |
| MOTA | 602 | CG1 | ILE | 66 | 60.332 | 50.062 | 14.586 | 1.00 14.56 | A_13 |
| MOTA | 603 | CD1 | ILE | 66 | 59.587 | 51.149 | 15.333 | 1.00 16.94 | A_13 |
| MOTA | 604 | C | ILE | 66 | 60.662 | 47.030 | 14.361 | 1.00 10.81 | A_13 |
| MOTA | 605 | 0 | ILE | 66 | 60.311 | 46.962 | 13.188 | 1.00 10.00 | A_13 |
| ATOM | 606 | N | LYS | 67 | 60.143 | 46.271 | 15.330 | 1.00 10.00 | A_13 |
| MOTA | 608 | CA | LYS | 67 | 59.036 | 45.327 | 15.103 | 1.00 10.23 | A_13 |
| MOTA | 609 | CB | LYS | 67 67 | 57.689 | 46.042 | 15.268 | 1.00 10.29 | A_13 |
| MOTA | 610 | CG | LYS | 67 67 | 57.584 | 46.895 | 16.510 | 1.00 14.63 | A_13 |
| MOTA MOTA | 611 612 | CE | LYS LYS | 67 67 | 57.646 | 46.056 | 17.774 | 1.00 14.94 | A_13 |
| ATOM | 613 | NZ | LYS | 67 67 | 57.382 57.480 | 46.923 | 18.986 | 1.00 22.99 | A_13 |
| MOTA | 617 | C | LYS | 67 | 59.113 | 46.174 44.633 | 20.258 13.726 | 1.00 28.27 | A_13 |
| MOTA | 618 | ŏ | LYS | 67 | 60.167 | | 13.726 | 1.00 17.91 1.00 24.16 | A_13 |
| ATOM | 619 | N | GLU | 68 | 58.027 | 44.690 | 12.949 | 1.00 12.72 | A_13 A_13 |
| ATOM | 621 | CA | GLU | 68 | 57.960 | 44.067 | 11.624 | 1.00 16.06 | A_13 |
| MOTA | 622 | CB | GLU | 68 | 56.505 | 44.019 | 11.128 | 1.00 26.89 | A_13 |
| ATOM | 623 | CG | GLU | 68 | 55.566 | 43.258 | 12.087 | 1.00 36.97 | A_13 |
| MOTA | 624 | CD | GLU | 68 | 54.217 | 43.973 | 12.381 | 1.00 41.61 | A_13 |
| ATOM | 625 | OE1 | | 68 | 53.289 | 43.921 | 11.537 | 1.00 17.31 | A_13 |
| ATOM | 626 | OE2 | | 68 | 54.074 | 44.561 | 13.485 | 1.00 26.72 | A_13 |
| ATOM ATOM | 627 628 | 0 | GLU | 68 | 58.823 | 44.911 | 10.705 | 1.00 22.50 | A_13 |
| ATOM | 629 | Ŋ | GLU HIS | 68 69 | 58.587 59.848 | 46.093 | 10.532 | 1.00 20.64 | A_13 |
| ATOM | 631 | CA | HIS | | 60.732 | 44.315 45.102 | 10.120 9.283 | 1.00 16.43 1.00 13.69 | A_13 |
| MOTA | 632 | CB | HIS | 69 | 61.930 | 45.603 | 10.103 | 1.00 10.97 | A_13 A_13 |
| MOTA | 633 | CG | HIS | 69 | 62.786 | 44.502 | 10.643 | 1.00 24.02 | A_13 |
| MOTA | 634 | CD2 | HIS | | 63.873 | 43.876 | 10.133 | 1.00 10.00 | A_13 |
| MOTA | 635 | ND1 | HIS | 69 | 62.512 | 43.876 | 11.839 | 1.00 17.68 | A_13 |
| MOTA | 637 | | HIS | | 63.384 | 42.912 | 12.041 | 1.00 12.53 | A_13 |
| MOTA | 638 | | HIS | | 64.228 | 42.888 | 11.020 | 1.00 10.00 | A_13 |
| ATOM | 639 | C | HIS | 69 | 61.214 | 44.469 | 7.983 | 1.00 21.28 | A_13 |
| MOTA | 640 | 0 | HIS | 69 70 | 62.314 | 44.780 | 7.529 | 1.00 18.74 | A_13 |
| MOTA | 641 | N | GLY | | 60.451 | | 7.411 | 1.00 13.11 | A_13 |
| ATOM ATOM | 643 644 | CA C | GLY | | 60.832 | | 6.127 | 1.00 10.00 | A_13 |
| ATOM | 645 | ŏ | GLY | | 61.262 61.523 | 41.533 | 5.936 | 1.00 10.00 | A_13 |
| MOTA | 646 | N | ASP | | 61.412 | | 4.794 | 1.00 15.12 1.00 19.99 | A_13 |
| ATOM | 648 | CA | ASP | | 61.842 | | 6.862 | 1.00 19.99 | A_13 |
| ATOM | 649 | CB | ASP | | 63.332 | | | 1.00 19.99 | A_13 A_13 |
| ATOM | 650 | CG | ASP | | 63.672 | | | 1.00 10.00 | A_13 A_13 |
| MOTA | 651 | | ASP | | 64.846 | | | 1.00 13.38 | A_13 |
| MOTA | 652 | OD2 | ASP | 71 | 62.774 | | | 1.00 12.94 | A_13 |
| MOTA | 653 | C | ASP | 71 | 60.998 | | | | A_13 |
| MOTA | 654 | 0 | ASP | | 61.319 | 37.190 | 7.649 | 1.00 24.45 | A_13 |
| ATOM | 655 | N | PHE | | 59.946 | | | 1.00 14.15 | A_13 |
| ATOM | 657 | CA | PHE | | 59.040 | | | 1.00 10.00 | A_13 |
| MOTA MOTA | 658 659 | CB | PHE | | 58.410 | | | 1.00 10.00 | A_13 |
| ATOM | 660 | CG CD1 | PHE PHE | | 57.360 56.115 | | | 1.00 10.00 | _ |
| ATOM | 661 | | PHE | | 57.624 | | | 1.00 23.01 1.00 12.52 | A_13 A_13 |
| | | | | | 023 | 507 | 3.717 | 12.72 | 4-73 |

| ATOM | 662 | CE1 | PHE | 72 | 55.144 | 38.290 | 6.950 | 1.00 18.99 | A_13 |
|--------------|------------|------------|------------|------------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 663 | | PHE | 72 | 56.662 | 38.023 | 5.091 | 1.00 13.37 | A_13 |
| MOTA | 664 | | PHE | 72 | 55.420 | 38.413 | 5.576 | 1.00 22.50 | A_13 |
| MOTA | 665 | | PHE | 72 72 | 59.634 | 37.523 | 10.392 | 1.00 16.31 | A_13 |
| MOTA MOTA | 666 667 | | PHE TYR | 72 73 | 59.111 60.737 | 36.596 38.141 | 11.021 10.793 | 1.00 15.64 1.00 18.10 | A_13 |
| ATOM | 669 | | TYR | 73 | 61.407 | 37.827 | 12.046 | 1.00 14.01 | A_13 A_13 |
| MOTA | 670 | | TYR | 73 | 62.845 | 37.331 | 11.803 | 1.00 21.08 | A_13 |
| MOTA | 671 | | TYR | 73 | 62.915 | 35.965 | 11.138 | 1.00 22.48 | A_13 |
| MOTA | 672 | | TYR | 73 | 63.579 | 35.788 | 9.923 | 1.00 30.23 | A_13 |
| ATOM | .673 | | TYR | 73 | 63.615 | 34.538 | 9.291 | 1.00 24.04 | A_13 |
| MOTA MOTA | 674 675 | | TYR TYR | 73 73 | 62.288 62.320 | 34.856 | 11.710 | 1.00 19.23 | A_13 |
| ATOM | 676 | | TYR | 73 | 62.984 | 33.606 33.460 | 11.083 9.875 | 1.00 29.35 1.00 12.50 | A_13 A_13 |
| ATOM | 677 | | TYR | 73 | 63.018 | 32.246 | 9.241 | 1.00 17.89 | A_13 |
| ATOM | 679 | | TYR | 73 | 61.360 | 39.203 | 12.721 | 1.00 22.00 | A_13 |
| MOTA | 680 | | TYR | 73 | 62.365 | 39.919 | 12.819 | 1.00 10.93 | A_13 |
| MOTA | 681 | N | PRO | 74 | 60.175 | 39.570 | 13.221 | 1.00 19.94 | A_13 |
| MOTA MOTA | 682 683 | CD CA | PRO PRO | 74 74 | 58.969 59.934 | 38.723 | 13.278 | 1.00 15.69 | A_13 |
| ATOM | 684 | CB | PRO | 7 4 | 58.417 | 40.843 | 13.886 14.067 | 1.00 16.75 1.00 17.27 | A_13 A_13 |
| MOTA | 685 | CG | PRO | 74 | 58.131 | 39.407 | 14.335 | 1.00 16.24 | A_13 |
| MOTA | 686 | C | PRO | 74 | 60.640 | 41.037 | 15.216 | 1.00 17.39 | A_13 |
| ATOM | 687 | 0 | PRO | 74 | 60.779 | 40.105 | 16.023 | 1.00 10.00 | A_13 |
| MOTA | 688 | N | PHE | 75 75 | 61.098 | 42.264 | 15.431 | 1.00 10.00 | A_13 |
| ATOM ATOM | 690 691 | CA CB | PHE | 75 75 | 61.743 62.613 | 42.618 43.865 | 16.675 | 1.00 16.45 | A_13 |
| MOTA | 692 | CG | PHE | 75 75 | 63.931 | 43.590 | 16.512 15.841 | 1.00 20.71 1.00 23.32 | A_13 A_13 |
| ATOM | 693 | | PHE | 75 | 64.694 | 42.482 | 16.200 | 1.00 12.03 | A_13 |
| MOTA | 694 | CD2 | | 75 | 64.405 | 44.420 | 14.842 | 1.00 22.30 | A_13 |
| MOTA | 695 | CE1 | | 75 | 65.905 | 42.214 | 15.572 | 1.00 17.64 | A_13 |
| MOTA | 696 | | PHE | 75 75 | 65.622 | 44.148 | 14.208 | 1.00 15.43 | A_13 |
| MOTA MOTA | 697 698 | CZ C | PHE | 75 75 | 66.367 60.632 | 43.044 | 14,576 | 1.00 10.00 | A_13 |
| ATOM | 699 | Ö | PHE | 75 75 | 59.443 | 42.784 42.778 | 17.707 17.370 | 1.00 25.73 1.00 18.57 | A_13 |
| ATOM | 700 | N | ASP | 76 | 61.009 | 43.002 | 18.952 | 1.00 20.50 | A_13 A_13 |
| MOTA | 702 | CA | ASP | 76 | 60.023 | 43.049 | 20.006 | 1.00 13.89 | A_13 |
| MOTA | 703 | CB | ASP | 76 | 60.241 | 41.805 | 20.873 | 1.00 20.69 | A_13 |
| ATOM | 704 | CG | ASP | 76 76 | 61.672 | 41.685 | 21.378 | 1.00 22.52 | A_13 |
| MOTA MOTA | 705 706 | ODI ODI | ASP | 76 76 | 61.947 | 40.771 | 22.174 | 1.00 20.06 | A_13 |
| ATOM | 707 | C | ASP | 76. 76 | 62.525 59.971 | 42.506 44.277 | 20.998 | 1.00 10.69 1.00 25.20 | A_13 |
| ATOM | 708 | Ö | ASP | 76 | 59.397 | 44.207 | 21.986 | 1.00 29.52 | A_13 A_13 |
| MOTA | 709 | N | GLY | 77 | 60.585 | 45.379 | 20.488 | 1.00 10.00 | A_13 |
| ATOM | 711 | CA | GLY | 77 | 60.575 | 46.553 | 21.334 | 1.00 10.00 | A_13 |
| MOTA | 712 | C | GLY | 77 | 61.769 | 46.514 | 22.266 | 1.00 10.00 | A_13 |
| ATOM ATOM | 713 | O N | GLY PRO | 77 78 | 62.735 61.785 | 45.797 47.344 | 21.987 | 1.00 18.49 | A_13 |
| MOTA | 715 | CD | PRO | 78 | 60.790 | 48.426 | 23.322 23.505 | 1.00 16.07 1.00 15.88 | A_13 A_13 |
| MOTA | 716 | CA | PRO | 78 | 62.855 | 47.439 | 24.330 | 1.00 16.23 | A_13 |
| MOTA | 717 | CB | PRO | 78 | 62.261 | 48.391 | 25.363 | 1.00 22.96 | A_13 |
| MOTA | 718 | CG | PRO | 78 70 | 61.470 | 49.349 | 24.501 | 1.00 22.37 | A_13 |
| MOTA MOTA | 719 720 | С 0 | PRO PRO | 78 78 | 63.150 | 46.090 | 24.969 | 1.00 25.32 | A_13 |
| MOTA | 721 | N | SER | 78 79 | 62.227 64.432 | 45.356 45.750 | 25.272 25.099 | 1.00 20.04 1.00 20.93 | A_13 |
| MOTA | 723 | CA | SER | 79 | 64.878 | 44.478 | 25.689 | 1.00 20.51 | A_13 A_13 |
| MOTA | 724 | CB | SER | 79 | 64.364 | 44.311 | 27.131 | 1.00 23.69 | A_13 |
| MOTA | 725 | OG | SER | 79 | 65.028 | 45.211 | 28.006 | 1.00 33.37 | A_13 |
| MOTA | 727 | C | SER | 79 70 | 64.557 | 43.248 | 24.863 | 1.00 20.39 | A_13 |
| MOTA MOTA | 728 729 | O N | SER GLY | 79 80 | 64.124 64.825 | 43.362 42.071 | 23.708 25.415 | 1.00 17.27 | A_13 |
| MOTA | 731 | CA | GLY | 80 | 64.564 | 40.850 | 24.678 | 1.00 13.38 1.00 10.11 | A_13 |
| MOTA | 732 | C | GLY | 80 | 65.471 | 40.808 | 23.458 | 1.00 13.15 | A_13 A_13 |
| MOTA | 733 | 0 | GLY | 80 | 66.614 | 41.251 | 23.538 | 1.00 31.80 | A_13 |
| MOTA | 734 | N | LEU | 81 | 64.939 | 40.393 | 22.310 | 1.00 29.05 | |
| MOTA | 736 | CA | LEU | 81 | 65.720 | 40.317 | 21.078 | 1.00 29.63 | A_13 |
| MOTA MOTA | 737 738 | CB CG | LEU | 81 81 | 64.789 | 40.033 | 19.905 | 1.00 19.67 | A_13 |
| ATOM | 739 | | LEU | 81 | 65.121 64.215 | 38.872 38.980 | 18.971 17.773 | 1.00 21.79 | A_13 |
| ATOM | 740 | | LEU | 81 | 66.590 | 38.918 | 18.518 | 1.00 23.87 1.00 22.09 | A_13 A_13 |
| MOTA | 741 | C | LEU | 81 | 66.442 | | | 1.00 19.25 | A_13 |
| MOTA | 742 | 0 | LEU | 81 | 65.808 | 42.700 | 20.872 | 1.00 14.95 | A_13 |
| ATOM | 743 | N | LEU | 82 | 67.760 | 41.599 | 20.657 | 1.00 25.03 | A_13 |
| MOTA MOTA | 745 746 | CA CB | LEU | 82 82 | 68.573 | 42.795 | 20.421 | 1.00 27.35 | A_13 |
| MOTA | 747 | CG | LEU | 82 82 | 69.868 69.802 | 42.747 42.748 | 21.244 22.773 | 1.00 12.74 | A_13 |
| ATOM | 748 | | LEU | 82 | 68.590 | 43.520 | 23.263 | 1.00 16.50 1.00 17.99 | A_13 A_13 |
| | | | _ | | | | | | 27 7 |

| ATOM 750 C LEU 82 68.938 42.945 18.949 1.00 24.79 | ATOM | 749 | CD2 | LEU | 82 | 69.744 | 41.343 | 23.279 | 1.00 13.28 | A_13 |
|--|------|-----|-----|-----|----|--------|--------|--------|------------|-------------|
| ATOM 752 N ALA 83 69,387 41,839 18,359 1.00 21,15 64 A_13 ATOM 755 CB ALA 83 69,780 41,819 16,961 1.00 15.64 A_13 ATOM 755 CB ALA 83 71,180 42,410 16,820 1.00 15.74 A_13 ATOM 755 C ALA 83 69,864 39,488 17,227 1.00 20,422 A_13 ATOM 755 N ALA 83 69,864 39,488 17,227 1.00 20,422 A_13 ATOM 758 N ALA 84 69,764 60,239 15,122 1.00 10,72 A_13 ATOM 758 N ALA 84 69,764 60,239 15,122 1.00 10,72 A_13 ATOM 758 N ALA 84 69,764 60,239 15,122 1.00 10,72 A_13 ATOM 761 CB ALS 84 69,764 60,239 15,122 1.00 10,72 A_13 ATOM 762 CG HIS 84 67,381 39,489 13,679 1.00 22,78 A_13 ATOM 763 CT2 HIS 84 67,381 39,489 13,679 1.00 22,78 A_13 ATOM 763 CT2 HIS 84 67,381 39,489 13,679 1.00 22,78 A_13 ATOM 766 CEL HIS 84 66,507 39,886 12,220 1.00 15,00 A_13 ATOM 769 O HIS 84 70,418 39,088 12,220 1.00 15,00 A_13 ATOM 769 O HIS 84 70,418 39,088 12,220 1.00 15,00 A_13 ATOM 769 O HIS 84 70,418 39,088 12,220 1.00 15,00 A_13 ATOM 770 N ALA 85 71,066 39,027 12,682 1.00 13,43 ATOM 770 N ALA 85 71,066 39,027 12,682 1.00 13,43 ATOM 770 N ALA 85 71,066 39,027 12,682 1.00 13,43 ATOM 770 N ALA 85 71,466 38,027 12,682 1.00 13,43 ATOM 770 N ALA 85 71,466 38,027 12,682 1.00 13,43 ALA ATOM 775 O ALA 85 71,466 36,000 ALA ATOM 775 O ALA 85 71,466 36,000 ALA ATOM 775 O ALA 85 70,448 ATOM 770 N ALA 86 70,448 ATOM 770 N ALA 870 ATOM 770 N ALA 86 70,448 ATOM 770 N ALA 870 ATOM 770 N ALA 870 ATOM 770 N ALA 870 ATOM 770 N ALA 88 ATOM 770 N ALA | | | | | | | | | | A_13 |
| ATOM 754 CA ALA 83 69,790 41,819 16,961 1.00 15.64 A_13 ATOM 755 CB ALA 83 71,180 42,410 16,820 1.00 15.74 A_13 ATOM 757 O ALA 83 69,806 40,400 16,001 15.74 A_13 ATOM 757 O ALA 83 69,806 40,400 16,001 15.74 A_13 ATOM 757 O ALA 83 69,806 40,400 15,126 1.00 10,123 A_13 ATOM 758 O ALA 83 69,806 40,400 15,126 1.00 10,72 A_13 ATOM 758 O CHIS 84 69,836 89,399 14,502 1.00 20,51 A_13 ATOM 760 C ALS 84 66,436 18,808 38,399 14,502 1.00 20,51 A_13 ATOM 761 CB ALS 84 66,436 18,808 38,399 14,502 1.00 20,51 A_13 ATOM 761 CB ALS 84 66,436 18,808 38,399 12,502 1.00 20,51 A_13 ATOM 766 CB ALS 84 66,436 18,808 38,399 12,502 1.00 20,51 A_13 ATOM 766 CB ALS 84 66,436 18,808 38,399 12,502 1.00 20,51 A_13 ATOM 766 CB ALS 84 66,502 18,869 14,104 1.00 13,50 A_13 ATOM 766 CB ALS 84 66,032 18,869 14,104 1.00 13,50 A_13 ATOM 766 CB ALS 84 66,032 18,869 14,104 1.00 13,50 A_13 ATOM 766 CB ALS 84 66,037 39,886 12,220 1.00 15,00 A_13 ATOM 769 O HIS 84 70,338 40,162 12,532 1.00 10,00 A_13 ATOM 769 O HIS 84 70,338 40,162 12,532 1.00 10,00 A_13 ATOM 770 CB ALS 85 71,746 37,983 11,402 1.00 10,00 A_13 ATOM 773 CB ALS 85 71,746 37,983 11,402 1.00 10,00 A_13 ATOM 775 CB ALS 85 71,746 37,983 11,402 1.00 10,00 A_13 ATOM 776 N PHE 86 71,459 15,565 11,402 1.00 10,00 A_13 ATOM 777 CB ALS 85 71,426 36,661 1,722 1.00 17,89 A_13 ATOM 777 CB ALS 85 71,426 36,661 1,722 1.00 17,89 A_13 ATOM 778 CB ALS 85 71,426 36,661 1,722 1.00 17,89 A_13 ATOM 779 CB PHE 86 67,149 13,7530 8,003 1.00 10,00 A_13 ATOM 779 CB PHE 86 67,149 13,7530 8,003 1.00 10,00 A_13 ATOM 779 CB PHE 86 67,149 13,7530 8,003 1.00 10,00 A_13 ATOM 779 CB PHE 86 67,129 17,134 7,13 | | | | | | | | | | |
| ATOM 755 CB ALA 83 71.180 (2.410 16.820 1.00 15.74 A.] ATOM 756 C ALA 83 69.866 40.00 16.444 1.00 19.37 A.] ATOM 757 O ALA 83 69.866 40.00 16.444 1.00 19.37 A.] ATOM 758 N HIS 84 69.746 40.252 15.126 1.00 10.72 A.] ATOM 758 N HIS 84 69.746 40.252 15.126 1.00 10.72 A.] ATOM 758 O ALA 83 69.866 38.939 14.502 1.00 10.72 A.] ATOM 758 O ALS 84 69.868 38.939 14.502 1.00 10.51 A.] ATOM 761 CD HIS 84 69.868 38.939 14.502 1.00 10.51 A.] ATOM 762 CD HIS 84 67.351 39.483 12.488 1.00 10.00 1.37 A.] ATOM 763 CD2 HIS 84 67.351 39.489 14.104 1.00 11.50 A.] ATOM 763 CD2 HIS 84 67.351 39.489 14.104 1.00 11.50 A.] ATOM 766 CEL HIS 84 65.307 39.497 13.210 1.00 10.00 A.] ATOM 767 NEZ HIS 84 65.307 39.497 13.210 1.00 14.37 A.] ATOM 768 C HIS 84 70.418 39.088 13.130 1.00 22.78 A.] ATOM 769 O HIS 84 70.418 39.088 13.130 1.00 22.78 A.] ATOM 770 NA ALA 85 71.086 38.027 12.685 1.00 13.43 A.] ATOM 770 NA ALA 85 71.086 38.027 12.685 1.00 10.00 A.] ATOM 771 C ALA 85 71.265 38.1027 12.685 1.00 10.00 A.] ATOM 772 C ALA 85 71.265 38.1027 12.685 1.00 10.00 A.] ATOM 773 C ALA 85 71.265 38.1027 12.685 1.00 10.00 A.] ATOM 775 C ALA 85 71.265 38.1027 12.685 1.00 10.00 A.] ATOM 775 C ALA 85 71.265 38.1027 12.685 1.00 10.00 A.] ATOM 775 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 778 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 778 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 778 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 778 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 779 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 779 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 779 C ALA 85 71.265 38.1027 12.685 1.00 12.49 A.] ATOM 780 C G PHE 86 71.697 38.502 8.002 1.00 10.00 A.] ATOM 781 C D PHE 86 71.697 38.502 8.002 1.00 10.00 A.] ATOM 780 C G PHE 86 71.697 38.002 8 | | | | | | | | | | - |
| ATOM 756 C ALA 83 69.806 40.400 16.444 1.00 19.37 | | | | | | | | | | |
| ATOM 757 O ALA 83 69.864 39.458 17.227 1.00 20.42 A_33 ATOM 760 CA HIS 84 69.808 38.939 14.502 1.00 20.51 A_13 ATOM 761 CB HIS 84 69.808 38.939 14.502 1.00 20.51 A_13 ATOM 761 CB HIS 84 69.808 38.939 14.502 1.00 20.51 A_13 ATOM 763 CD HIS 84 67.361 38.849 13.679 10.00 24.79 A_13 ATOM 763 CD HIS 84 67.361 38.849 13.679 10.00 24.79 A_13 ATOM 763 CD HIS 84 67.361 38.849 13.679 10.00 24.79 A_13 ATOM 766 CD HIS 84 67.361 38.849 11.679 10.00 24.79 A_13 ATOM 766 CD HIS 84 66.052 38.869 14.104 1.00 13.50 A_13 ATOM 766 CD HIS 84 66.052 38.869 14.104 1.00 13.50 A_13 ATOM 768 CD HIS 84 66.052 38.869 14.104 1.00 13.50 A_13 ATOM 768 CD HIS 84 60.083 39.888 12.220 1.00 10.00 A_13 ATOM 768 CD HIS 84 60.083 39.888 12.220 1.00 10.00 A_13 ATOM 770 N ALA 85 71.086 38.027 12.885 1.00 13.43 AL3 ATOM 770 N ALA 85 71.086 38.027 12.885 1.00 13.43 AL3 ATOM 773 CB ALA 85 71.242 36.661 10.721 1.00 17.89 A_13 ATOM 774 C ALA 85 71.426 36.661 10.721 1.00 17.89 A_13 ATOM 775 O ALA 85 70.900 35.746 11.346 1.00 19.43 A_13 ATOM 776 N PHE 86 71.459 35.372 8.651 1.00 12.49 A_13 ATOM 776 CB PHE 86 71.697 36.6585 9.425 1.00 13.49 A_13 ATOM 776 CB PHE 86 70.739 35.728 7.344 1.00 10.00 A_13 ATOM 778 CB PHE 86 70.739 35.728 7.344 1.00 10.00 A_13 ATOM 778 CB PHE 86 70.739 35.728 7.344 1.00 10.00 A_13 ATOM 778 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 778 CB PHE 86 67.739 35.728 7.344 1.00 10.00 A_13 ATOM 778 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 778 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 778 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A_13 ATOM 789 CB PHE 86 69.432 36.240 7.529 1.00 19.96 A | | | | | | | | | | |
| ATOM 760 CA HIS 84 69.808 38.939 14.502 1.00 20.51 Å_13 ATOM 761 CS HIS 84 67.361 38.849 13.679 10.0 24.79 Å_13 ATOM 763 CD2 HIS 84 67.361 38.849 13.679 10.0 24.79 Å_13 ATOM 763 CD2 HIS 84 67.361 39.849 12.268 1.00 10.00 Å_13 ATOM 766 CD1 HIS 84 66.052 38.869 14.104 1.00 13.50 Å_13 ATOM 766 CD1 HIS 84 66.052 38.869 14.104 1.00 13.50 Å_13 ATOM 766 CD1 HIS 84 66.087 39.886 12.220 1.00 15.00 Å_13 ATOM 768 C HIS 84 70.318 84.092 12.251 1.00 10.00 Å_13 ATOM 769 O HIS 84 70.318 40.162 12.532 1.00 10.00 Å_13 ATOM 769 O HIS 84 70.318 40.162 12.532 1.00 10.00 Å_13 ATOM 779 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 773 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 773 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 775 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 775 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 775 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 775 CB ALA 85 71.746 37.983 11.402 1.00 10.03 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 775 CB ALA 85 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 778 CB A PHE 86 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 778 CB A PHE 86 71.697 36.585 9.45 1.00 13.49 Å_13 ATOM 778 CB A PHE 86 71.697 36.585 9.40 1.00 16.63 Å_13 ATOM 780 CB PHE 86 71.697 36.585 9.40 1.00 16.63 Å_13 ATOM 780 CB PHE 86 69.148 36.240 7.529 1.00 19.966 Å_13 ATOM 780 CB PHE 86 69.149 37.530 8.003 1.00 10.63 Å_13 ATOM 780 CB PHE 86 69.149 37.530 8.003 1.00 10.00 Å_13 ATOM 780 CB PHE 86 69.149 37.530 8.003 1.00 10.00 Å_13 ATOM 780 CB PHE 86 69.598 37.298 38.009 8.158 1.00 19.06 Å_13 ATOM 780 CB PHE 86 69.149 37.398 38.009 8.158 1 | | | | | | | | | | |
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| ATOM 817 CB ASN 91 68.863 36.932 -0.999 1.00 15.26 A 13 ATOM 818 CG ASN 91 68.860 36.430 -2.439 1.00 36.74 A 13 ATOM 819 OD1 ASN 91 68.497 35.282 -2.701 1.00 29.56 A 13 ATOM 820 ND2 ASN 91 69.265 37.286 -3.376 1.00 27.03 A 13 ATOM 823 C ASN 91 70.226 37.986 0.849 1.00 24.66 A 13 ATOM 824 O ASN 91 71.257 38.479 1.313 1.00 17.43 A 13 ATOM 825 N TYR 92 69.198 37.632 1.622 1.00 17.69 A 13 ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A 13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A 13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A 13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A 13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A 13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A 13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A 13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A 13 | | | | | | | | | | |
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| ATOM 819 OD1 ASN 91 68.497 35.282 -2.701 1.00 29.56 A 13 ATOM 820 ND2 ASN 91 69.265 37.286 -3.376 1.00 27.03 A 13 ATOM 823 C ASN 91 70.226 37.986 0.849 1.00 24.66 A 13 ATOM 824 O ASN 91 71.257 38.479 1.313 1.00 17.43 A 13 ATOM 825 N TYR 92 69.198 37.632 1.622 1.00 17.69 A 13 ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A 13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A 13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A 13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A 13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A 13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A 13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A 13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A 13 | | | | | | | | | | |
| ATOM 820 ND2 ASN 91 69.265 37.286 -3.376 1.00 27.03 A_13 ATOM 823 C ASN 91 70.226 37.986 0.849 1.00 24.66 A_13 ATOM 824 O ASN 91 71.257 38.479 1.313 1.00 17.43 A_13 ATOM 825 N TYR 92 69.198 37.632 1.622 1.00 17.69 A_13 ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A_13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A_13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A_13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | | | | | | |
| ATOM 823 C ASN 91 70.226 37.986 0.849 1.00 24.66 A_13 ATOM 824 O ASN 91 71.257 38.479 1.313 1.00 17.43 A_13 ATOM 825 N TYR 92 69.198 37.632 1.622 1.00 17.69 A_13 ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A_13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A_13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A_13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | 91 | | 37.286 | -3.376 | 1.00 27.03 | A_13 |
| ATOM 825 N TYR 92 69.198 37.632 1.622 1.00 17.69 A 13 ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A 13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A 13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A 13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A 13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A 13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A 13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A 13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A 13 | | | | | | | | | | A_13 |
| ATOM 827 CA TYR 92 69.233 37.876 3.061 1.00 10.17 A_13 ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A_13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A_13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | | | | | | |
| ATOM 828 CB TYR 92 67.942 37.428 3.744 1.00 16.78 A_13 ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A_13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | | | | | | |
| ATOM 829 CG TYR 92 66.786 38.364 3.523 1.00 26.17 A_13 ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | • | | | | | | |
| ATOM 830 CD1 TYR 92 66.015 38.803 4.581 1.00 17.79 A_13 ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | MOTA | 829 | CG | TYR | 92 | 66.786 | 38.364 | | | |
| ATOM 831 CE1 TYR 92 64.947 39.678 4.380 1.00 29.60 A_13 ATOM 832 CD2 TYR 92 66.467 38.818 2.250 1.00 25.90 A_13 ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A_13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | 92 | | | 4.581 | 1.00 17.79 | A_13 |
| ATOM 833 CE2 TYR 92 65.406 39.691 2.040 1.00 30.60 A 13 ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A 13 | | | | | | | | | | |
| ATOM 834 CZ TYR 92 64.647 40.117 3.107 1.00 12.31 A_13 | | | | | | | | | | |
| Amous And a | | | | | | | | | | |
| | | | | | | | | | | |

| ATOM | 837 | c ' | TYR | 92 | 70.427 | 37.245 | 3.763 | 1.00 11.94 | A_13 |
|--------------|------------|----------|------------|------------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 838 | | TYR | 92 | 70.752 | 37.617 | 4.882 | 1.00 17.58 | A_13 |
| ATOM | 839 | N (| GLY | 93 | 71.095 | 36.311 | 3.097 | 1.00 24.67 | A_13 |
| ATOM | 841 | CA | GLY | 93 | 72.250 | 35.666 | 3.691 | 1.00 18.05 | A_13 |
| ATOM | 842 | _ | GLY | 93 | 73.295 | 36.681 | 4.116 | 1.00 10.00 | A_13 |
| ATOM | 843 | | GLY | 93 | 73.573 | 37.656 | 3.391 | 1.00 10.13 | A_13 |
| MOTA | 844 | - | GLY | 94 | 73.812 | 36.495 | 5.328 | 1.00 12.44 | A_13 |
| ATOM | 846 | | GLY | 94 | 74.827 | 37.372 | 5.872 | 1.00 10.00 | A_13 |
| MOTA | 847 | | GLY | 94 | 74.358 | 38.694 | 6.456 | 1.00 17.29 | A_13 |
| ATOM | 848 | _ | GLY | 94 | 75.052 | 39.271 | 7.284 | 1.00 14.53 | A_13 |
| MOTA | 849 | | ASP | 95 | 73.221 | 39.206 | 5.993 | 1.00 10.00 | A_13 |
| MOTA MOTA | 851 852 | | ASP ASP | 95 95 | 72.689 | 40.485 | 6.472 | 1.00 16.35 | A_13 |
| ATOM | 853 | | ASP | 95 | 71.332 71.421 | 40.777 40.904 | 5.814 | 1.00 10.00 1.00 14.54 | A_13 |
| MOTA | 854 | | ASP | 95 | 70.406 | 41.256 | 4.309 3.673 | 1.00 14.54 | A_13 A_13 |
| ATOM | 855 | | ASP | 95 | 72.502 | 40.647 | 3.753 | 1.00 15.39 | A_13 A_13 |
| MOTA | 856 | | ASP | 95 | 72.548 | 40.523 | 7.994 | 1.00 22.31 | A_13 |
| ATOM | 857 | | ASP | 95 | 72.279 | 39.497 | 8.635 | 1.00 10.88 | A_13 |
| ATOM | 858 | | ALA | 96 | 72.703 | 41.711 | 8.566 | 1.00 18.45 | A_13 |
| MOTA | 860 | ÇA | ALA | 96 | 72.609 | 41.877 | 10.011 | 1.00 15.08 | A_13 |
| ATOM | 861 | | ALA | 96 | 73.982 | 42.244 | 10.587 | 1.00 19.20 | A_13 |
| ATOM | 862 | | ALA | 96 | 71.587 | 42.961 | 10.345 | 1.00 14.91 | A_13 |
| ATOM | 863 | | ALA | 96 | 71.702 | 44.092 | 9.876 | 1.00 10.00 | A_13 |
| ATOM | 864 | | HIS | 97 | 70.635 | 42.646 | 11.215 | 1.00 14.01 | A_13 |
| MOTA | 866 | | HIS | . 97 | 69.599 | 43.620 | 11.581 | 1.00 11.35 | A_13 |
| MOTA MOTA | 867 868 | | HIS HIS | 97 97 | 68.207 | 43.083 | 11.203 | 1.00 20.32 | A_13 |
| ATOM | 869 | | HIS | 97 | 68.027 68.734 | 42.786 43.186 | 9.742 | 1.00 15.00 1.00 10.00 | A_13 |
| MOTA | 870 | ND1 | | 97 | 67.014 | 41.978 | 8.654 9.257 | 1.00 10.00 | A_13 |
| MOTA | 871 | CEI | | 97 | 67.108 | 41.895 | 7.936 | 1.00 10.00 | A_13 A_13 |
| ATOM | 872 | NE2 | | 97 | 68.142 | 42.618 | 7.552 | 1.00 17.10 | A_13 |
| MOTA | 874 | C | HIS | 97 | 69.650 | 43.952 | 13.078 | 1.00 13.37 | A_13 |
| ATOM | 875 | | HIS | 97 | | . 43.055 | 13.908 | 1.00 13.48 | A_13 |
| MOTA | 876 | N | PHE | 98 | 69.596 | 45.237 | 13.423 | 1.00 21.01 | A_13 |
| MOTA | 878 | CA | PHE | 98 | 69.634 | 45.668 | 14.823 | 1.00 11.27 | A_13 |
| MOTA | 879 | CB | PHE | 98 | 70.817 | 46.615 | 15.055 | 1.00 10.00 | A_13 |
| MOTA | 880 | CG | PHE | 98 | 72.138 | 46.011 | 14.703 | 1.00 20.49 | A_13 |
| ATOM | 881 | CD1 | PHE | 98 | 72.984 | 45.524 | 15.707 | 1.00 17.49 | A_13 |
| ATOM | 882 | CD2 | PHE | 98 | 72.506 | 45.853 | 13.365 | 1.00 13.51 | A_13 |
| MOTA | 883 | CE1 | PHE | 98 | 74.171 | 44.888 | 15.382 | 1.00 20.00 | A_13 |
| MOTA | 884 | CE2 | PHE | 98 | 73.693 | 45.215 | 13.024 | 1.00 10.00 | A_13 |
| ATOM ATOM | 885 | CZ | PHE | 98 | 74.527 | 44.728 | 14.029 | 1.00 10.00 | A_13 |
| MOTA | 886 887 | CO | PHE PHE | 98 98 | 68.336 67.815 | 46.336 | 15.245 | 1.00 25.38 | A_13 |
| ATOM | 888 | N | ASP | 99 | 67.817 | 47.218 45.924 | 14.552 16.394 | 1.00 10.00 1.00 21.68 | A_13 |
| ATOM | 890 | CA | ASP | 99 | 66.567 | 46.476 | 16.886 | 1.00 21.00 | A_13 A_13 |
| ATOM | 891 | CB | ASP | 99 | 66.039 | 45.604 | 18.010 | 1.00 10.00 | A_13 |
| ATOM | 892 | CG | ASP | 99 | 64.648 | 45.998 | 18.473 | 1.00 14.00 | A_13 |
| MOTA | 893 | OD1 | ASP | 99 | 64.104 | 45.272 | 19.329 | 1.00 15.19 | A_13 |
| MOTA | 894 | OD2 | ASP | 99 | 64.089 | 47.011 | 18.001 | 1.00 17.01 | A_13 |
| MOTA | 895 | C | ASP | 99 | 66.817 | 47.871 | 17.391 | 1.00 13.06 | A_13 |
| ATOM | 896 | 0 | ASP | 99 | 67.528 | 48.056 | 18.374 | 1.00 10.00 | A_13 |
| ATOM | 897 | N | ASP | 100 | 66.203 | 48.856 | 16.746 | 1.00 15.56 | A_13 |
| ATOM | 899 | CA | ASP | 100 | 66.397 | 50.232 | 17.177 | 1.00 18.23 | A_13 |
| MOTA MOTA | 900 901 | CB CG | ASP ASP | 100 100 | 66.121 | 51.228 | 16.041 | 1.00 15.05 | |
| MOTA | 902 | | ASP | 100 | 67.275 67.602 | 52.180 52.516 | 15.838 | 1.00 11.67 | A_13 |
| ATOM | 903 | | ASP | 100 | 67.879 | 52.569 | 14.683 16.860 | 1.00 21.07 1.00 14.72 | A_13 |
| ATOM | 904 | C | ASP | 100 | 65.610 | 50.572 | 18.445 | 1.00 10.00 | A_13 A_13 |
| MOTA | 905 | Ö | ASP | 100 | 65.767 | 51.635 | 19.009 | 1.00 17.18 | A_13 |
| MOTA | 906 | N | ASP | 101 | 64.755 | | 18.895 | 1.00 14.57 | A_13 |
| MOTA | 908 | CA | ASP | 101 | 64.031 | 49.924 | 20.123 | 1.00 17.59 | A_13 |
| MOTA | 909 | CB | ASP | 101 | 62.769 | 49.051 | 20.236 | 1.00 12.50 | A_13 |
| MOTA | 910 | CG | ASP | 101 | 61.532 | 49.721 | 19.606 | 1.00 17.12 | A_13 |
| ATOM | 911 | | | 101 | 60.599 | 49.023 | 19.179 | 1.00 10.39 | A_13 |
| MOTA | 912 | OD2 | | 101 | 61.480 | | 19.536 | 1.00 18.09 | A_13 |
| MOTA | 913 | C | ASP | 101 | 64.994 | 49.766 | 21.306 | 1.00 19.33 | A_13 |
| MOTA | 914 | 0 | ASP | 101 | 64.610 | | 22.456 | 1.00 10.00 | A_13 |
| MOTA | 915 | N | GLU | 102 | 66.213 | 49.301 | 21.019 | 1.00 16.15 | A_13 |
| ATOM | 917 | CA | GLU | 102 | 67.267 | | 22.044 | 1.00 13.43 | A_13 |
| MOTA MOTA | 918 919 | CB | GLU | 102 | 68.264 | | 21.720 | | A_13 |
| MOTA | 920 | CG CD | GLU GLU | 102 102 | 67.697 66.650 | | 21.636 | 1.00 10.00 | A_13 |
| ATOM | 921 | OE1 | | 102 | 66.872 | | 22.672 23.870 | 1.00 11.18 1.00 16.09 | A_13 |
| MOTA | 922 | OE2 | | 102 | 65.572 | | 22.271 | 1.00 16.09 | A_13 A_13 |
| ATOM | 923 | c | GLU | 102 | 68.070 | | 22.007 | 1.00 20.70 | A_13 |
| ATOM | 924 | 0 | GLU | 102 | 68.103 | | 20.971 | 1.00 13.97 | A_13 |
| | | | | | | • | | | |

| ATOM | 925 | N : | THR | 103 | 68.774 | 50.823 | 23.091 | 1.00 22.82 | A_13 |
|--------------|--------------|----------|------------|------------|------------------|------------------|------------------|--------------------------|---------------|
| ATOM ATOM | 927 928 | | THR THR | 103 103 | 69.606 69.571 | 52.034 52.793 | 23.102 24.459 | 1.00 13.45 1.00 20.78 | A_13 A_13 |
| MOTA | 929 | OG1 ' | THR | 103 | 68.236 | 53.228 | 24.745 | 1.00 10.69 | A_13 |
| MOTA ATOM | 931 932 | | THR THR | 103 103 | 70.445 71.030 | 54.046 51.571 | 24.378 22.822 | 1.00 19.45 1.00 12.42 | A_13 A_13 |
| MOTA | 933 | 0 | THR | 103 | 71.639 | 50.896 | 23.642 | 1.00 19.81 | A_13 |
| ATOM ATOM | 934 936 | | TRP TRP | 104 104 | 71.525 72.873 | 51.854 51.448 | 21.626 21.248 | 1.00 10.00 1.00 13.61 | A_13 A_13 |
| ATOM | 937 | CB | TRP | 104 | 72.943 | 51.221 | 19.739 | 1.00 29.21 | A_13 |
| ATOM ATOM | 938 939 | | TRP TRP | 104 104 | 71.970 72.101 | 50.174 48.760 | 19.313 19.501 | 1.00 21.39 1.00 25.13 | A_13 A_13 |
| MOTA | 940 | CE2 | TRP | 104 | 70.937 | 48.156 | 18.964 | 1.00 28.84 | A_13 A_13 |
| MOTA ATOM | 941 942 | | TRP TRP | 104 104 | 73.088 70.765 | 47.941 50.372 | 20.070 18.694 | 1.00 13.36 1.00 21.59 | A_13 |
| MOTA | 943 | NE1 | TRP | 104 | 70.139 | 49.163 | 18.484 | 1.00 19.91 | A_13 A_13 |
| MOTA MOTA | 945 946 | | TRP TRP | 104 104 | 70.738 72.888 | 46.768 46.568 | 18.977 20.084 | 1.00 10.00 1.00 14.54 | A_13 A_13 |
| MOTA | 947 | CH2 | TRP | 104 | 71.720 | 45.995 | 19.539 | 1.00 11.93 | A_13 |
| ATOM ATOM | 948 949 | C | TRP TRP | 104 104 | 73.912 73.707 | 52.453 53.671 | 21.725 21.642 | 1.00 16.59 1.00 12.90 | A_13 |
| ATOM | 950 | N | THR | 105 | 75.013 | 51.949 | 22.268 | 1.00 20.85 | A_13 A_13 |
| ATOM ATOM | 952 953 | CA CB | THR THR | 105 105 | 76.040 75.974 | 52.831 52.890 | 22.794 24.322 | 1.00 12.38 1.00 14.39 | A_13 |
| ATOM . | 954 | OG1 | THR | 105 | 76.345 | 51.609 | 24.849 | 1.00 14.39 | A_13 A_13 |
| ATOM ATOM | 956 957 | CG2 C | THR THR | 105 105 | 74.575 77.437 | 53.273 52.378 | 24.797 22.457 | 1.00 12.17 1.00 10.00 | A_13 |
| MOTA | 958 | 0 | THR | 105 | 77.644 | 51.261 | 22.012 | 1.00 18.98 | A_13 A_13 |
| MOTA MOTA | 959 961 | N CA | SER SER | 106 106 | 78.385 79.809 | 53.277 53.043 | 22.704 22.502 | 1.00 26.01 1.00 17.80 | A_13 |
| MOTA | 962 | CB | SER | 106 | 80.466 | 54.284 | 21.888 | 1.00 17.80 | A_13 A_13 |
| MOTA MOTA | 963 965 | OG C | SER SER | 106 106 | 79.744 80.435 | 54.756 52.779 | 20.763 23.880 | 1.00 38.89 1.00 34.75 | A_13 |
| MOTA | 966 | Ö | SER | 106 | 81.652 | 52.884 | 24.042 | 1.00 34.75 | A_13 A_13 |
| ATOM ATOM | 967 969 | N CA | SER SER | 107 107 | 79.590 80.032 | 52.494 52.221 | 24.875 26.240 | 1.00 25.87 1.00 19.68 | A_13 |
| ATOM | 970 | CB | SER | 107 | 80.082 | 53.510 | 27.061 | 1.00 19.88 | A_13 A_13 |
| ATOM ATOM | 971 973 | OG C | SER SER | 107 107 | 78.819 79.100 | 54.158 | 27.096 | 1.00 33.70 | A_13 |
| MOTA | 974 | Ö | SER | 107 | 78.460 | 51.200 50.418 | 26.892 26.193 | 1.00 13.60 1.00 16.40 | A_13 A_13 |
| ATOM ATOM | 975 977 | N CA | SER SER | 108 108 | 79.028 78.188 | 51.205 | 28.221 | 1.00 17.31 | A_13 |
| ATOM | 978 | CB | SER | 108 | 78.745 | 50.259 50.009 | 28.949 30.364 | 1.00 20.12 1.00 22.63 | A_13 A_13 |
| ATOM ATOM | 979 981 | OG C | SER SER | 108 108 | 78.444 76.702 | 51.061 50.606 | 31.271 29.076 | 1.00 27.69 1.00 19.98 | A_13 |
| ATOM | 982 | Ö | SER | 108 | 75.921 | 49.785 | 29.562 | 1.00 15.96 | A_13 A_13 |
| MOTA MOTA | 983 985 | N CA | LYS LYS | 109 109 | 76.311 74.907 | 51.820 52.186 | 28.713 28.847 | 1.00 16.24 1.00 11.10 | · A_13 |
| MOTA | 986 | CB | LYS | 109 | 74.740 | 53.688 | 28.690 | 1.00 12.41 | A_13 A_13 |
| MOTA MOTA | 987 988 | CG CD | LYS LYS | 109 109 | 73.555 | 54.239 55.732 | 29.462 29.258 | 1.00 32.67 1.00 25.94 | A_13 |
| MOTA | 989 | CE | LYS | 109 | 74.535 | 56.599 | 29.749 | 1.00 25.34 | A_13 A_13 |
| ATOM ATOM | 990 994 | NZ C | LYS LYS | 109 109 | 74.225 74.138 | 58.070 | 29.636 27.773 | 1.00 22.70 1.00 21.67 | A_13 A_13 |
| MOTA | 995 | 0 | LYS | 109 | 74.667 | 51.210 | 26.694 | 1.00 32.76 | A_13 |
| MOTA MOTA | 996 998 | N CA | GLY GLY | 110 110 | 72.932 72.156 | 50.955 50.206 | 28.081 27.096 | 1.00 29.60 1.00 10.31 | A_13 A_13 |
| MOTA | 999 | C | GLY | 110 | 72.965 | 49.043 | 26.542 | 1.00 20.08 | A_13 |
| ATOM ATOM | 1000 1001 | O N | GLY TYR | 110 111 | 73.672 72.924 | 48.362 48.859 | 27.285 25.227 | 1.00 11.17 1.00 12.05 | A_13 A_13 |
| MOTA | 1003 | CA | TYR | 111 | 73.665 | 47.791 | 24.583 | 1.00 13.45 | A_13 |
| MOTA MOTA | 1004 1005 | CB CG | TYR TYR | 111 111 | 72.713 71.776 | 46.871 46.101 | 23.806 24.716 | 1.00 21.16 1.00 12.28 | A_13 A_13 |
| MOTA | 1006 | CD1 | TYR | 111 | 70.455 | 46.510 | 24.906 | 1.00 14.85 | A_13 |
| MOTA MOTA | 1007 1008 | CE1 | TYR TYR | 111 111 | 69.618 72.232 | 45.837 44.995 | 25.795 25.435 | 1.00 19.08 1.00 21.86 | A_13 A_13 |
| MOTA | 1009 | CE2 | TYR | 111 | 71.405 | 44.314 | 26.324 | 1.00 10.00 | A_13 |
| MOTA MOTA | 1010 1011 | CZ OH | TYR TYR | 111 111 | 70.101 69.282 | 44.740 | | 1.00 18.51 1.00 14.32 | A_13 .A_13 |
| MOTA | 1013 | C | TYR | 111 | 74.779 | 48.335 | 23.695 | 1.00 16.73 | A_13 |
| MOTA MOTA | 1014 1015 | N O | TYR ASN | 111 112 | 74.540 76.008 | | | 1.00 11.98 1.00 11.80 | A_13 A_13 |
| ATOM ATOM | 1017 1018 | CA | ASN | 112 | 77.184 | 48.357 | 23.240 | 1.00 16.37 | A_13 |
| ATOM | 1018 | CB CG | asn asn | 112 112 | 78.453 79.701 | 47.867 48.460 | | 1.00 27.52 1.00 20.16 | A_13 A_13 |
| MOTA MOTA | 1020 1021 | OD1 | ASN | 112 | 80.327 | 47.861 | 22.447 | 1.00 20.99 | A_13 |
| al VII | 1021 | MD2 | ASN | 112 | 80.082 | 49.640 | 23.801 | 1.00 15.12 | A_13 |

| 3.000 | 1004 | _ | | | | | | | |
|-------|------|-----|-----|-------|----------|--------|--------|------------|--------|
| MOTA | 1024 | C | ASN | 112 | 77.137 | 47.809 | 21.813 | 1.00 18.08 | A_13 |
| ATOM | 1025 | 0 | ASN | 112 | 77.288 | 46.606 | 21.592 | 1.00 12.69 | A_13 |
| MOTA | 1026 | N | LEU | 113 | 76.972 | 48.700 | 20.844 | 1.00 11.15 | A_13 |
| ATOM | 1028 | CA | LEU | 113 | 76.878 | 48.296 | 19.461 | 1.00 10.00 | A_13 |
| ATOM | 1029 | CB | LEU | 113 | 76.718 | 49.526 | 18.568 | 1.00 10.24 | A_13 |
| ATOM | 1030 | CG | LEU | 113 | 76.325 | 49.262 | 17.106 | 1.00 15.67 | |
| MOTA | 1031 | CD1 | LEU | 113 | 75.155 | | | | A_13 |
| | | | | | | 48.296 | 17.050 | 1.00 26.54 | A_13 |
| MOTA | 1032 | CD2 | LEU | 113 | 75.967 | 50.555 | 16.415 | 1.00 15.60 | A_13 |
| MOTA | 1033 | C | LEU | 113 | 78.037 | 47.403 | 18.986 | 1.00 25.17 | A_13 |
| MOTA | 1034 | 0 | LEU | 113 | 77.799 | 46.380 | 18.336 | 1.00 17.24 | A_13 |
| ATOM | 1035 | N | PHE | 114 | 79.274 | 47.759 | 19.327 | 1.00 28.89 | _ |
| ATOM | 1037 | CA | PHE | 114 | 80.442 | 46.974 | | | A_13 |
| | | | | | | | 18.910 | 1.00 19.15 | A_13 |
| ATOM | 1038 | CB | PHE | 114 | 81.753 | 47.579 | 19.434 | 1.00 14.60 | A_13 |
| MOTA | 1039 | CG | PHE | 114 | 82.923 | 46.627 | 19.374 | 1.00 18.53 | A_13 |
| MOTA | 1040 | CD1 | PHE | 114 | 83.419 | 46.175 | 18.144 | 1.00 26.13 | A_13 |
| MOTA | 1041 | CD2 | PHE | 114 | 83.514 | 46.162 | 20.547 | 1.00 17.22 | A_13 |
| MOTA | 1042 | CE1 | PHE | 114 | 84.475 | 45.271 | 18.086 | 1.00 10.43 | A_13 |
| MOTA | 1043 | CE2 | PHE | 114 | 84.571 | 45.259 | 20.502 | 1.00 16.51 | |
| ATOM | 1044 | CZ | | | | | | | A_13 |
| | | | PHE | 114 | 85.052 | 44.815 | 19.260 | 1.00 15.54 | A_13 |
| MOTA | 1045 | С | PHE | 114 | 80.359 | 45.508 | 19.306 | 1.00 10.00 | A_13 |
| MOTA | 1046 | 0 | PHE | 114 | 80.437 | 44.625 | 18.445 | 1.00 33.07 | A_13 |
| ATOM | 1047 | N | LEU | 115 | 80.206 | 45.249 | 20.600 | 1.00 12.18 | A_13 |
| ATOM | 1049 | CA | LEU | 115 | 80.113 | 43.877 | 21.103 | 1.00 10.59 | A_13 |
| ATOM | 1050 | CB | LEU | 115 | 79.874 | 43.895 | 22.616 | 1.00 14.14 | A_13 |
| MOTA | 1051 | CG | LEU | 115 | 81.082 | 43.937 | 23.578 | | |
| ATOM | 1052 | | | | | | | 1.00 34.39 | A_13 |
| | | | LEU | 115 | 82.337 | 44.354 | 22.863 | 1.00 14.93 | A_13 |
| ATOM | 1053 | CD2 | LEU | 115 | 80.815 | 44.836 | 24.793 | 1.00 13.42 | A_13 |
| MOTA | 1054 | C | LEU | 115 | 79.019 | 43.080 | 20.379 | 1.00 12.06 | A_13 |
| MOTA | 1055 | 0 | LEU | 115 | 79.298 | 42.109 | 19.675 | 1.00 13.35 | A_13 |
| MOTA | 1056 | N | VAL | 116 | 77.786 | 43.558 | 20.459 | 1.00 13.11 | A_13 |
| MOTA | 1058 | CA | VAL | 116 | 76.678 | 42.875 | 19.814 | 1.00 12.97 | |
| ATOM | 1059 | CB | VAL | 116 | 75.343 | | | | A_13 |
| | | | | | | 43.569 | 20.129 | 1.00 28.07 | A_13 |
| MOTA | 1060 | | VAL | 116 | 74.200 | | 19.340 | 1.00 17.32 | A_13 |
| MOTA | 1061 | | VAL | 116 | 75.074 | 43.491 | 21.617 | 1.00 22.14 | A_13 |
| ATOM | 1062 | C | VAL | 116 | 76.862 | 42.724 | 18.313 | 1.00 10.00 | A_13 |
| ATOM | 1063 | 0 | VAL | 116 | 76.473 | 41.716 | 17.755 | 1.00 14.68 | A_13 |
| MOTA | 1064 | N | ALA | 117 | 77.481 | 43.706 | 17.667 | 1.00 10.80 | |
| ATOM | 1066 | CA | ALA | 117 | 77.726 | | | | A_13 |
| ATOM | 1067 | | | | | 43.662 | 16.224 | 1.00 18.28 | A_13 |
| | | CB | ALA | 117 | 78.223 | 45.014 | 15.727 | 1.00 14.94 | A_13 |
| MOTA | 1068 | С | ALA | 117 | 78.735 | 42.579 | 15.863 | 1.00 25.24 | A_13 |
| MOTA | 1069 | 0 | ALA | 117 | 78.562 | 41.872 | 14.861 | 1.00 18.50 | A_13 |
| ATOM | 1070 | N | ALA | 118 | 79.795 | 42.458 | 16.665 | 1.00 24.40 | A_13 |
| ATOM | 1072 | CA | ALA | 118 | 80.829 | 41.451 | 16.422 | 1.00 11.80 | A_13 |
| ATOM | 1073 | CB | ALA | 118 | 81.945 | 41.590 | | _ | |
| MOTA | 1074 | C | | | | | 17.447 | 1.00 19.28 | A_13 |
| | | | ALA | 118 | 80.178 | 40.056 | 16.496 | 1.00 10.00 | A_13 |
| ATOM | 1075 | 0 | ALA | 118 | 80.426 | 39.183 | 15.660 | 1.00 10.00 | A_13 |
| ATOM | 1076 | N | HIS | 119 | 79.309 | 39.875 | 17.487 | 1.00 19.01 | A_13 |
| MOTA | 1078 | CA | HIS | 119 | 78.587 | 38.624 | 17.674 | 1.00 14.36 | A_13 |
| MOTA | 1079 | CB | HIS | 119 | 77.725 | 38.751 | 18.924 | 1.00 10.00 | A_13 |
| ATOM | 1080 | CG | HIS | 119 | • 76.796 | 37.602 | 19.166 | 1.00 10.00 | A_13 |
| ATOM | 1081 | CD2 | | 119 | 75.691 | 37.187 | 18.498 | 1.00 14.94 | |
| MOTA | 1082 | | HIS | 119 | 76.905 | | | | A_13 |
| ATOM | 1084 | | | | | 36.783 | 20.263 | 1.00 20.37 | A_13 |
| | | | HIS | 119 | 75.917 | 35.909 | 20.270 | 1.00 17.53 | A_13 |
| MOTA | 1085 | | HIS | 119 | 75.161 | 36.134 | | 1.00 17.55 | A_13 |
| MOTA | 1086 | С | HIS | 119 | 77.741 | 38.339 | 16.419 | 1.00 10.00 | A_13 |
| ATOM | 1087 | 0 | HIS | 119 | 77.779 | 37.245 | 15.856 | 1.00 10.64 | A_13 |
| ATOM | 1088 | N | GLŲ | 120 | 77.004 | 39.343 | 15.968 | 1.00 22.95 | A_13 |
| MOTA | 1090 | CA | GLÜ | 120 | 76.174 | 39.224 | 14.775 | 1.00 23.96 | A_13 |
| MOTA | 1091 | CB | GLU | 120 . | 75.429 | 40.545 | 14.502 | 1.00 17.19 | |
| ATOM | 1092 | CG | GLU | 120 | 74.373 | | | | A_13 |
| ATOM | 1093 | | | | | 40.889 | 15.555 | 1.00 16.14 | A_13 |
| | | CD | GLU | 120 | 73.492 | 39.691 | 15.929 | 1.00 10.00 | A_13 |
| ATOM | 1094 | OE1 | | 120 | 73.478 | 39.354 | 17.122 | 1.00 17.94 | A_13 |
| MOTA | 1095 | OE2 | GLU | 120 | 72.844 | 39.078 | 15.047 | 1.00 17.03 | A_13 |
| MOTA | 1096 | C | GLU | 120 | 76.992 | 38.832 | | 1.00 11.45 | A_13 |
| ATOM | 1097 | 0 | GLU | 120 | 76.594 | 37.946 | 12.772 | 1.00 13.34 | A_13 |
| ATOM | 1098 | N | PHE | 121 | 78.127 | 39.498 | | 1.00 10.00 | 2 13 |
| ATOM | 1100 | CA | PHE | | | | | | A_13 |
| | | | | 121 | 78.959 | | | 1.00 14.70 | A_13 |
| MOTA | 1101 | CB | PHE | 121 | 80.040 | 40.245 | | 1.00 10.00 | A_13 |
| ATOM | 1102 | CG | PHE | 121 | 79.481 | 41.623 | 11.792 | 1.00 21.57 | A_13 |
| MOTA | 1103 | CD1 | PHE | 121 | 80.235 | 42.764 | | 1.00 16.73 | A_13 |
| MOTA | 1104 | CD2 | PHE | 121 | 78.164 | | | 1.00 13.91 | · A_13 |
| MOTA | 1105 | | PHE | 121 | 79.682 | 44.054 | 11.891 | 1.00 11.69 | |
| ATOM | 1106 | CE2 | | 121 | 77.615 | 43.066 | | | A_13 |
| ATOM | 1107 | | | | | | | 1.00 18.93 | A_13 |
| | | CZ | PHE | 121 | 78.373 | 44.192 | | 1.00 10.00 | A_13 |
| MOTA | 1108 | C | PHE | 121 | 79.505 | 37.756 | | 1.00 17.14 | A_13 |
| ATOM | 1109 | 0 | PHE | 121 | 79.642 | 37.104 | | 1.00 13.04 | A_13 |
| ATOM | 1110 | N | GLY | 122 | 79.738 | 37.245 | 13.490 | 1.00 16.60 | A_13 |
| | | | | | | | | | |

| | | | | | • | | | | |
|------|------|-----|-----|-----|----------|--------|--------|------------|--------|
| ATOM | 1112 | CA | GLY | 122 | 80.202 | 35.872 | 13.627 | 1.00 19.45 | n 12 |
| ATOM | 1113 | C | GLY | 122 | | | | | A_13 |
| | | | | | 79.162 | 34.982 | 12.966 | 1.00 18.55 | A_13 |
| ATOM | 1114 | 0 | GLY | 122 | 79.500 | 33.988 | 12.306 | 1.00 10.03 | A_13 |
| ATOM | 1115 | N | HIS | 123 | 77.892 | 35.361 | 13.140 | 1.00 18.22 | A_13 |
| ATOM | 1117 | CA | HIS | 123 | 76.753 | 34.665 | 12.525 | 1.00 16.31 | A_13 |
| MOTA | 1118 | CB | HIS | 123 | 75.424 | 35.224 | 13.031 | 1.00 11.35 | A_13 |
| MOTA | 1119 | CG | HIS | 123 | 75.049 | 34.768 | 14.403 | 1.00 10.33 | A_13 |
| MOTA | 1120 | CD2 | HIS | 123 | 74.552 | 35.454 | 15.457 | 1.00 16.64 | A_13 |
| MOTA | 1121 | ND1 | HIS | 123 | . 75.097 | 33.450 | 14.782 | 1.00 18.04 | A_13 |
| MOTA | 1123 | CE1 | HIS | 123 | 74.638 | 33.332 | 16.017 | 1.00 16.66 | A_13 |
| MOTA | 1124 | NE2 | HIS | 123 | 74.301 | 34.533 | 16.450 | 1.00 25.32 | A_13 |
| MOTA | 1125 | C | HIS | 123 | 76.771 | 34.853 | 10.997 | 1.00 13.66 | |
| MOTA | 1126 | Õ | HIS | 123 | 76.565 | 33.901 | | | A_13 |
| | | | | | | | 10.246 | 1.00 10.82 | A_13 |
| MOTA | 1127 | N | SER | 124 | 77.006 | 36.082 | 10.539 | 1.00 13.57 | A_13 |
| MOTA | 1129 | CA | SER | 124 | 77.030 | 36.368 | 9.099 | 1.00 12.03 | A_13 |
| MOTA | 1130 | CB | SER | 124 | 77.311 | 37.863 | 8.832 | 1.00 10.35 | A_13 |
| MOTA | 1131 | OG | SER | 124 | 76.399 | 38.706 | 9.510 | 1.00 14.26 | A_13 |
| MOTA | 1133 | С | SER | 124 | 78.117 | 35.548 | 8.422 | 1.00 21.45 | A_13 |
| MOTA | 1134 | 0 | SER | 124 | 78.079 | 35.333 | 7.210 | 1.00 10.00 | A_13 |
| MOTA | 1135 | N | LEU | 125 | 79.091 | 35.108 | 9.216 | 1.00 10.00 | A_13 |
| MOTA | 1137 | CA | LEU | 125 | 80.222 | 34.340 | 8.707 | 1.00 19.28 | A_13 |
| MOTA | 1138 | CB | LEU | 125 | 81.521 | 34.754 | 9.422 | 1.00 22.39 | A_13 |
| MOTA | 1139 | CG | LEU | 125 | 81.849 | 36.258 | 9.340 | 1.00 10.00 | A_13 |
| ATOM | 1140 | CD1 | | 125 | 83.063 | 36.622 | 10.190 | 1.00 10.00 | A_13 |
| MOTA | 1141 | CD2 | LEU | 125 | 82.029 | 36.651 | 7.873 | 1.00 10.00 | |
| ATOM | 1142 | C | LEU | 125 | 79.986 | 32.851 | | | A_13 |
| MOTA | 1143 | Ö | | 125 | | | 8.843 | 1.00 10.00 | A_13 |
| | | | LEU | | 80.759 | 32.056 | 8.329 | 1.00 23.27 | A_13 |
| MOTA | 1144 | N | GLY | 126 | 78.932 | 32.477 | 9.563 | 1.00 22.87 | A_13 |
| MOTA | 1146 | CA | GLY | 126 | 78.604 | 31.070 | 9.720 | 1.00 17.27 | A_13 |
| MOTA | 1147 | С | GLY | 126 | 78.781 | 30.464 | 11.094 | 1.00 11.71 | A_13 |
| MOTA | 1148 | 0 | GLY | 126 | 78.784 | 29.244 | 11.236 | 1.00 24.16 | A_13 |
| MOTA | 1149 | N | LEU | 127 | 78.972 | 31.297 | 12.105 | 1.00 18.95 | A_13 |
| MOTA | 1151 | CA | LEU | 127 | 79.152 | 30.790 | 13.457 | 1.00 22.84 | A_13 |
| MOTA | 1152 | CB | LEU | 127 | 80.113 | 31.693 | 14.252 | 1.00 11.92 | A_13 |
| MOTA | 1153 | CG | LEU | 127 | 81.244 | 30.969 | | 1.00 18.83 | A_13 |
| MOTA | 1154 | | LEU | 127 | 82.096 | 30.197 | 13.979 | 1.00 16.63 | |
| MOTA | 1155 | CD2 | | 127 | 82.104 | | | _ | A_13 |
| MOTA | 1156 | C | LEU | 127 | | 31.970 | 15.760 | 1.00 22.15 | A_13 |
| | | | | | 77.802 | 30.699 | 14.163 | 1.00 21.02 | A_13 |
| MOTA | 1157 | 0 | LEU | 127 | 76.996 | 31.629 | 14.098 | 1.00 14.68 | A_13 |
| ATOM | 1158 | N | ASP | 128 | 77.563 | 29.572 | 14.828 | 1.00 18.87 | A_13 |
| MOTA | 1160 | CA | ASP | 128 | 76.336 | 29.345 | 15.571 | 1.00 16.46 | A_13 |
| ATOM | 1161 | CB | ASP | 128 | 75.996 | 27.855 | 15.540 | 1.00 17.60 | A_13 |
| MOTA | 1162 | CG | ASP | 128 | 74.577 | 27.552 | 15.996 | 1.00 23.55 | A_13 |
| MOTA | 1163 | OD1 | ASP | 128 | 73.796 | 28.488 | 16.258 | 1.00 10.00 | A_13 |
| MOTA | 1164 | OD2 | ASP | 128 | 74.236 | 26.355 | 16.087 | 1.00 32.36 | A_13 |
| MOTA | 1165 | C | ASP | 128 | 76.634 | 29.803 | 16.995 | 1.00 10.00 | A_13 |
| ATOM | 1166 | Ö | ASP | 128 | 77.650 | 30.420 | 17.244 | 1.00 29.54 | A_13 |
| ATOM | 1167 | N | HIS | 129 | 75.714 | 29.565 | 17.912 | 1.00 10.00 | |
| ATOM | 1169 | CA | HIS | 129 | 75.910 | 29.955 | | | A_13 |
| ATOM | 1170 | CB | HIS | 129 | | | 19.289 | 1.00 10.00 | A_13 |
| ATOM | | | | | 74.582 | 30.033 | 20.029 | 1.00 21.30 | A_13 |
| | 1171 | CG | HIS | 129 | 73.798 | 31.282 | 19.761 | 1.00 24.16 | A_13 |
| ATOM | 1172 | CD2 | | 129 | 74.180 | 32.585 | 19.725 | 1.00 10.00 | A_13 |
| ATOM | 1173 | | HIS | 129 | 72.460 | 31.263 | 19.476 | 1.00 21.70 | A_13 |
| ATOM | 1175 | | HIS | 129 | 72.031 | 32.501 | 19.271 | 1.00 10.27 | A_13 |
| ATOM | 1176 | NE2 | | 129 | 73.057 | 33.319 | 19.407 | 1.00 14.37 | A_13 |
| MOTA | 1177 | Ç | HIS | 129 | 76.780 | 28.947 | 19.992 | 1.00 30.04 | A_13 |
| MOTA | 1178 | 0 | HIS | 129 | 76.624 | 27.730 | 19.822 | 1.00 22.13 | A_13 |
| MOTA | 1179 | N | SER | 130 | 77.628 | 29.468 | 20.860 | 1.00 18.60 | A_13 |
| ATOM | 1181 | CA | SER | 130 | 78.534 | 28.662 | 21.636 | 1.00 10.79 | . A_13 |
| ATOM | 1182 | CB | SER | | 79.849 | 29.435 | | 1.00 21.31 | A_13 |
| MOTA | 1183 | OG | SER | | 80.782 | 28.731 | | | |
| ATOM | 1185 | C | SER | | | | | | A_13 |
| ATOM | 1186 | | | | 77.898 | 28.368 | | 1.00 31.13 | A_13 |
| | | 0 | SER | | 76.962 | 29.060 | | 1.00 15.87 | A_13 |
| ATOM | 1187 | N | LYS | | 78.402 | 27.319 | | 1.00 13.13 | A_13 |
| ATOM | 1189 | CA | LYS | | 77.924 | 26.925 | | | A_13 |
| ATOM | 1190 | CB | LYS | | 77.656 | 25.414 | 24.990 | 1.00 18.85 | A_13 |
| MOTA | 1191 | CG | LYS | 131 | 78.689 | 24.541 | | 1.00 32.55 | A_13 |
| ATOM | 1192 | CD | LYS | 131 | 78.547 | 24.601 | | | A_13 |
| ATOM | 1193 | CE | LYS | | 79.909 | 24.672 | | | A_13 |
| ATOM | 1194 | NZ | LYS | | 80.747 | 25.799 | | | |
| ATOM | 1198 | C | LYS | | 78.922 | 27.379 | | | A_13 |
| ATOM | 1199 | Ö | LYS | | 78.666 | 27.260 | | | A_13 |
| ATOM | 1200 | N | ASP | | | | | | A_13 |
| ATOM | 1202 | | | | 80.025 | 27.968 | | | A_13 |
| ATOM | | CA | ASP | | 81.097 | 28.487 | | | A_13 |
| | 1203 | CB | ASP | | 82.376 | 28.617 | | | A_13 |
| ATOM | 1204 | CG | ASP | | 83.649 | 28.821 | | | A_13 |
| MOTA | 1205 | נמט | ASP | 132 | 84.645 | 28.132 | 26.028 | 1.00 36.08 | A_13 |

| ATOM | 1206 | OD2 | ASP | 132 | 83.685 | 29.660 | 27.276 | 1.00 15.60 | A_13 |
|--------------|--------------|------------|------------|--------------|------------------|------------------|------------------|--------------------------|--------------|
| MOTA MOTA | 1207 1208 | С 0 | ASP ASP | 132 132 | 80.603 80.559 | 29.875 30.816 | 26.836 | 1.00 18.74 | A_13 |
| MOTA | 1209 | N | PRO | 133 | 80.305 | 30.039 | 26.038 28.142 | 1.00 14.61 1.00 15.61 | A_13 A_13 |
| ATOM | 1210 | CD | PRO | 133 | 80.617 | 29.127 | 29.251 | 1.00 21.19 | A_13 |
| ATOM ATOM | 1211 1212 | CA CB | PRO PRO | 133 133 | 79.818 79.542 | 31.320 | 28.662 | 1.00 10.00 | A_13 |
| ATOM | 1213 | CG | PRO | 133 | 80.633 | 31.007 30.063 | 30.135 30.450 | 1.00 10.00 1.00 30.94 | A_13 A_13 |
| MOTA | 1214 | C | PRO | 133 | 80.834 | 32.444 | 28.511 | 1.00 22.87 | A_13 |
| ATOM | 1215 | 0 | PRO | 133 | 80.526 | 33.574 | 28.742 | 1.00 21.65 | A_13 |
| MOTA MOTA | 1216 1218 | N CA | GLY | 134 134 | 82.070 83.055 | 32.115 33.167 | 28.174 28.028 | 1.00 20.95 1.00 15.22 | A_13 |
| ATOM | 1219 | C | GLY | 134 | 83.182 | 33.578 | 26.581 | 1.00 15.22 1.00 34.54 | A_13 A_13 |
| MOTA | 1220 | 0 | GLY | 134 | 83.962 | 34.488 | 26.252 | 1.00 18.06 | A_13 |
| MOTA MOTA | 1221 1223 | N CA | ALA ALA | 135 135 | 82.490 82.547 | 32.846 | 25.706 | 1.00 21.09 | A_13 |
| ATOM | 1224 | CB | ALA | 135 | 82.131 | 33.110 31.858 | 24.263 23.453 | 1.00 27.50 1.00 10.00 | A_13 A_13 |
| ATOM | 1225 | C | ALA | 135 | 81.722 | 34.308 | 23.814 | 1.00 21.74 | A_13 |
| ATOM ATOM | 1226 1227 | N O | ALA LEU | 135 . 136 | 80.641 | 34.556 | 24.328 | 1.00 13.84 | A_13 |
| ATOM | 1229 | CA | LEU | 136 | 82.220 81.540 | 34.990 36.140 | 22.787 22.203 | 1.00 19.10 1.00 21.65 | A_13 |
| ATOM | 1230 | CB | LEU | 136 . | 82.448 | 36.803 | 21.161 | 1.00 10.00 | A_13 A_13 |
| ATOM ATOM | 1231 1232 | CG CD1 | LEU | 136 | 81.964 | 37.898 | 20.201 | 1.00 17.22 | A_13 |
| ATOM | 1233 | CD2 | | 136 136 | 81.250 81.113 | 37.296 38.896 | 19.024 20.905 | 1.00 24.18 1.00 10.00 | A_13 |
| ATOM | 1234 | C | LEU | 136 | 80.250 | 35.632 | 21.558 | 1.00 19.32 | A_13 A_13 |
| ATOM | 1235 | 0 | LEU | 136 | 79.266 | 36.359 | 21.458 | 1.00 26.20 | A_13 |
| MOTA MOTA | 1236 1238 | N CA | MET MET | 137 137 | 80.297 79.123 | 34.409 33.791 | 21.029 20.423 | 1.00 10.00 | A_13 |
| ATOM | 1239 | CB | MET | 137 | 79.507 | 32.691 | 19.428 | 1.00 10.02 1.00 15.14 | A_13 A_13 |
| MOTA | 1240 | CG | MET | 137 | 80.181 | 33.223 | 18.169 | 1.00 16.42 | A_13 |
| ATOM ATOM | 1241 1242 | SD CE | MET MET | 137 137 | 79.366 | 34.665 | 17.397 | 1.00 10.65 | A_13 |
| ATOM | 1243 | CE | MET | 137 | 77.848 78.122 | 34.005 33.256 | 16.975 21.447 | 1.00 10.87 1.00 12.70 | A_13 |
| MOTA | 1244 | 0 | MET | 137 | 77.187 | 32.539 | 21.087 | 1.00 10.00 | A_13 A_13 |
| ATOM | 1245 | N | PHE | 138 | 78.295 | 33.627 | 22.713 | 1.00 18.70 | A_13 |
| MOTA MOTA | 1247 1248 | CA CB | PHE | 138 138 | 77.370 77.954 | 33.196 33.448 | 23.759 | 1.00 24.08 | A_13 |
| ATOM | 1249 | CG | PHE | 138 | 77.306 | 32.617 | 25.159 26.240 | 1.00 24.15 1.00 29.38 | A_13 A_13 |
| MOTA | 1250 | CD1 | | 138 | 76.694 | 33.222 | 27.336 | 1.00 27.07 | A_13 |
| MOTA MOTA | 1251 1252 | CD2 CE1 | | 138 138 | 77.253 | 31.226 | 26.123 | 1.00 21.37 | A_13 |
| ATOM | 1253 | CE2 | | 138 | 76.033 76.599 | 32.455 30.458 | 28.289 27.065 | 1.00 30.35 1.00 19.58 | A_13 A_13 |
| ATOM | 1254 | CZ | PHE | 138 | 75.986 | 31.070 | 28.154 | 1.00 17.69 | A_13 |
| MOTA MOTA | 1255 1256 | C | PHE | 138 | 76.074 | 33.992 | 23.513 | 1.00 14.20 | A_13 |
| ATOM | 1257 | O N | PHE | 138 139 | 76.115 74.899 | 35.105 33.366 | 23.014 23.730 | 1.00 10.27 1.00 13.04 | A_13 |
| MOTA | 1258 | CD | PRO | 139 | 74.664 | 31.975 | | 1.00 13.04 | A_13 A_13 |
| MOTA | 1259 | CA | PRO | 139 | 73.619 | 34.043 | 23.504 | 1.00 18.27 | A_13 |
| MOTA MOTA | 1260 1261 | CB CG | PRO PRO | 139 139 | 72.625 73.474 | 32.875 31.634 | 23.384 | 1.00 14.33 | A_13 |
| MOTA | 1262 | C | PRO | 139 | 73.162 | 35.018 | 23.305 24.584 | 1.00 24.22 1.00 16.51 | A_13 A_13 |
| MOTA | 1263 | 0 | PRO | 139 | 72.023 | 35.467 | 24.535 | 1.00 24.45 | A_13 |
| MOTA MOTA | 1264 1266 | N CA | ILE | 140 140 | 74.034 | 35.375 | 25.524 | 1.00 23.16 | A_13 |
| MOTA | 1267 | CB | ILE | | 73.652 73.688 | 36.290 35.559 | 26.604 27.966 | 1.00 25.00 1.00 12.10 | A_13 A_13 |
| MOTA | 1268 | CG2 | | 140 | 73.336 | 36.519 | 29.085 | 1.00 12.62 | A_13 |
| ATOM ATOM | 1269 1270 | CG1 CD1 | | 140 140 | 72.738 | 34.341 | 27.904 | 1.00 22.67 | A_13 |
| ATOM | 1271 | C | ILE | 140 | 72.827 74.584 | 33.353 37.489 | 29.073 26.621 | 1.00 27.73 1.00 30.64 | A_13 |
| ATOM | 1272 | 0 | ILE | 140 | 75.778 | 37.317 | 26.682 | 1.00 30.04 | A_13 A_13 |
| MOTA | 1273 | N | TYR | | 74.033 | 38.694 | 26.532 | 1.00 21.05 | A_13 |
| MOTA MOTA | 1275 1276 | CA CB | TYR TYR | | 74.851 74.017 | 39.901 41.122 | 26.528 | 1.00 20.10 | A_13 |
| ATOM | 1277 | CG | TYR | | 74.784 | 42.433 | 26.129 26.103 | 1.00 17.66 1.00 22.24 | A_13 A_13 |
| MOTA | 1278 | CD1 | TYR | 141 | 74.711 | 43.318 | 27.171 | 1.00 18.07 | A_13 |
| ATOM ATOM | 1279 1280 | CE1 | TYR TYR | | 75.386 | 44.527 | 27.144 | 1.00 19.84 | A_13 |
| ATOM | 1281 | CE2 | | | 75.563 76.244 | 42.798 44.008 | 24.999 24.961 | 1.00 18.08 1.00 10.00 | A_13 |
| MOTA | 1282 | CZ | TYR | 141 | 76.149 | 44.867 | 26.038 | 1.00 10.00 | A_13 A_13 |
| ATOM ATOM | 1283 | ОН | TYR | 141 | 76.814 | 46.070 | 26.043 | 1.00 30.78 | A_13 |
| MOTA | 1285 1286 | C 0 . | TYR TYR | | 75.533 74.910 | 40.169 | 27.852 | 1.00 19.61 | A_13 |
| MOTA | 1287 | N | THR | 142 | 76.817 | 40.146 | 28.913 27.772 | 1.00 16.08 1.00 26.26 | A_13 A_13 |
| MOTA | 1289 | CA | THR | 142 | 77.612 | 40.788 | 28.944 | 1.00 24.52 | A_13 |
| MOTA MOTA | 1290 1291 | CB OG1 | THR THR | 142 | 78.498 | 39.568 | 29.362 | 1.00 10.00 | A_13 |
| | *671 | ΩI | ını | 142 | 77.664 | 38.587 | 29.981 | 1.00 37.30 | A_13 |

| MOTA | 1293 | CG2 | THR | 142 | 79.543 | 39.961 | 30.390 | 1.00 14.88 | A_13 |
|--------------|--------------|-----------|------------|------------|------------------|------------------|------------------|--------------------------|--------------|
| MOTA | 1294 | C | THR | 142 | 78.467 | 41.976 | 28.580 | 1.00 25.46 | A_13 |
| ATOM | 1295 | | THR | 142 | 78.980 | 42.058 | 27.464 | 1.00 10.00 | A_13 |
| ATOM ATOM | 1296 1298 | N CA | TYR TYR | 143 143 | 78.575 79.412 | 42.947 44.079 | 29.476 29.133 | 1.00 20.23 1.00 32.69 | A_13 |
| MOTA | 1299 | CB | TYR | 143 | 79.024 | 45.363 | 29.854 | 1.00 35.01 | A_13 A_13 |
| ATOM | 1300 | CG | TYR | 143 | 79.834 | 46.531 | 29.347 | 1.00 16.01 | A_13 |
| MOTA | 1301 | CD1 | TYR | 143 | 79.776 | 46.910 | 27.998 | 1.00 12.56 | A_13 |
| ATOM | 1302 | CE1 | TYR | 143 | 80.554 | 47.961 | 27.510 | 1.00 19.23 | A_13 |
| MOTA | 1303 | | TYR | 143 | 80.690 | 47.230 | 30.196 | 1.00 19.43 | A_13 |
| MOTA MOTA | 1304 1305 | CE2 | TYR TYR | 143 143 | 81.478 81.403 | 48.287 | 29.719 | 1.00 15.52 1.00 12.56 | A_13 |
| ATOM | 1305 | OH | TYR | 143 | 82.193 | 49.654 | 28.376 27.892 | 1.00 12.36 | A_13 A_13 |
| MOTA | 1308 | C | TYR | 143 | 80.871 | 43.754 | 29.382 | 1.00 25.10 | A_13 |
| MOTA | 1309 | 0 | TYR | 143 | 81.373 | 43.846 | 30.503 | 1.00 28.90 | A_13 |
| ATOM | 1310 | N | THR | 144 | 81.539 | 43.375 | 28.303 | 1.00 35.25 | A_13 |
| MOTA | 1312 | CA | THR | 144 | 82.946 | 43.029 | 28.336 | 1.00 38.86 | A_13 |
| ATOM ATOM | 1313 1314 | CB OG1 | THR THR | 144 144 | 83.158 82.129 | 41.568 41.219 | 27.873 26.934 | 1.00 23.22 1.00 35.22 | A_13 |
| MOTA | 1316 | CG2 | THR | 144 | 83.105 | 40.616 | 29.082 | 1.00 33.22 | A_13 A_13 |
| MOTA | 1317 | C | THR | 144 | 83.720 | 44.017 | 27.488 | 1.00 21.63 | A_13 |
| MOTA | 1318 | 0 | THR | 144 | 84.434 | 43.651 | 26.556 | 1.00 37.44 | A_13 |
| ATOM | 1319 | N | GLY | 145 | 83.504 | 45.288 | 27.798 | 1.00 14.47 | A_13 |
| MOTA | 1321 | CA | GLY | 145 | 84.200 | 46.375 | 27.131 | 1.00 24.39 | A_13 |
| ATOM ATOM | 1322 1323 | 0 | GLY GLY | 145 145 | 84.119 84.053 | 46.536 45.565 | 25.628 24.877 | 1.00 41.65 1.00 42.39 | A_13 |
| MOTA | 1324 | N | LYS | 146 | 84.122 | 47.792 | 25.195 | 1.00 42.39 | A_13 A_13 |
| MOTA | 1326 | CA | LYS | 146 | 84.059 | 48.103 | 23.778 | 1.00 29.29 | A_13 |
| MOTA | 1327 | CB | LYS | 146 | 83.260 | 49.392 | 23.539 | 1.00 26.47 | A_13 |
| MOTA | 1328 | CG | LYS | 146 | 83.087 | 49.721 | 22.059 | 1.00 33.24 | A_13 |
| MOTA | 1329 | CD | LYS | 146 | 82.812 | 51.194 | 21.833 | 1.00 13.70 | A_13 |
| ATOM ATOM | 1330 1331 | CE NZ | LYS LYS | 146 146 | 82.620 83.766 | 51.497 | 20.343 | 1.00 18.35 | A_13 |
| MOTA | 1335 | C | LYS | 146 | 85.491 | 51.122 48.297 | 19.477 23.308 | 1.00 30.66 1.00 41.61 | A_13 A_13 |
| ATOM | 1336 | ō | LYS | 146 | 86.028 | 49.412 | 23.382 | 1.00 46.44 | A_13 |
| ATOM | 1337 | N | SER | 147 | 86.130 | 47.206 | 22.898 | 1.00 34.67 | A_13 |
| MOTA | 1339 | CA | SER | 147 | 87.509 | 47.258 | 22.416 | 1.00 30.76 | A_13 |
| MOTA | 1340 | CB | SER | 147 | 87.624 | 48.258 | 21.249 | 1.00 24.56 | A_13 |
| ATOM | 1341 | OG | SER | 147 | 86.638 | 48.002 | 20.257 | 1.00 31.81 | A_13 |
| MOTA MOTA | 1343 1344 | C O | SER SER | 147 147 | 88.464 88.789 | 47.626 48.806 | 23.567 23.789 | 1.00 33.60 1.00 39.96 | A_13 |
| ATOM | 1345 | N | HIS | 148 | 88.862 | 46.611 | 24.331 | 1.00 35.36 | A_13 A_13 |
| MOTA | 1347 | CA | HIS | 148 | 89.778 | 46.769 | 25.467 | 1.00 34.40 | A_13 |
| MOTA | 1348 | CB | HIS | 148 | 89.307 | 47.862 | 26.438 | 1.00 26.40 | A_13 |
| ATOM | 1349 | CG | HIS | 148 | 90.251 | 49.022 | 26.537 | 1.00 39.11 | A_13 |
| MOTA MOTA | 1350 1351 | CD2 | HIS HIS | 148 148 | 90.929 | 49.542 | 27.588 | 1.00 30.52 | A_13 |
| ATOM | 1353 | | HIS | 148 | 90.635 91.511 | 49.767 50.681 | 25.437 25.807 | 1.00 37.71 1.00 29.04 | A_13 A_13 |
| ATOM | 1354 | NE2 | | 148 | 91.707 | 50.567 | 27.110 | 1.00 29.03 | A_13 |
| MOTA | 1356 | C | HIS | 148 | 89.949 | 45.436 | 26.190 | 1.00 39.41 | A_13 |
| ATOM | 1357 | 0 | HIS | 148 | 90.134 | 45.373 | 27.411 | 1.00 35.01 | A_13 |
| MOTA | 1358 | N | PHE | 149 | 89.840 | 44.386 | 25.383 | 1.00 25.35 | A_13 |
| MOTA MOTA | 1360 1361 | CA CB | PHE | 149 149 | 89.996 88.788 | 42.966 42.423 | 25.721 26.495 | 1.00 30.54 1.00 33.34 | A_13 A_13 |
| MOTA | 1362 | CG | PHE | 149 | 88.951 | 42.440 | 27.996 | 1.00 33.34 | A_13 |
| MOTA | 1363 | CD1 | PHE | 149 | 89.387 | 41.302 | 28.673 | 1.00 30.46 | A_13 |
| MOTA | 1364 | | PHE | 149 | 88.624 | 43.575 | 28.740 | 1.00 40.67 | A_13 |
| ATOM | 1365 | | PHE | 149 | 89.492 | 41.293 | 30.075 | 1.00 18.92 | A_13 |
| MOTA MOTA | 1366 1367 | CEZ | PHE | 149 149 | 88.728 | 43.574 | 30.136 | 1.00 23.23 | A_13 |
| MOTA | 1368 | C | PHE | 149 | 89.161 90.026 | 42.430 42.366 | 30.803 24.295 | 1.00 17.03 1.00 41.76 | A_13 A_13 |
| MOTA | 1369 | Õ | PHE | 149 | 89.967 | 43.119 | | 1.00 40.43 | A_13 |
| MOTA | 1370 | N | MET | 150 | 90.132 | 41.050 | 24.142 | 1.00 31.30 | A_13 |
| ATOM | 1372 | CA | MET | 150 | 90.152 | 40.531 | 22.779 | 1.00 20.65 | A_13 |
| MOTA | 1373 | CB | MET | 150 | 91.588 | _ | | 1.00 28.29 | A_13 |
| ATOM ATOM | 1374 1375 | CG | MET | 150 150 | 92.494 | 41.436 | 22.188 | 1.00 34.71 | A_13 |
| ATOM | 1375 | SD CE | MET MET | 150 150 | 91.750 92.512 | 42.780 42.498 | 21.185 | 1.00 67.91 | A_13 |
| ATOM | 1377 | C | MET | 150 | 89.201 | 39.370 | | 1.00 22.43 1.00 21.51 | A_13 A_13 |
| MOTA | 1378 | ŏ | MET | 150 | 88.498 | 38.901 | 23.391 | 1.00 25.37 | A_13 |
| ATOM | 1379 | N | LEU | 151 | 89.159 | 38.938 | 21.240 | | A_13 |
| MOTA | 1381 | CA | LEU | 151 | 88.313 | | 20.834 | 1.00 14.73 | A_13 |
| MOTA MOTA | 1382 | CB | LEU | 151 | 88.435 | 37.589 | | 1.00 15.49 | A_13 |
| ATOM | 1383 1384 | CG | LEU LEU | 151 151 | 87.535 86.070 | 36.511 36.915 | 18.691 | 1.00 27.05 | A_13 |
| ATOM | 1385 | | LEU | 151 | 87.879 | 36.310 | | 1.00 10.98 1.00 15.73 | A_13 A_13 |
| MOTA | 1386 | C | LEU | 151 | 88.732 | 36.563 | _ | | A_13 |
| | | | | | | | | | |

| MOTA | 1387 | 0 | LEU | 151 | 89.912 | 36.178 | 21.589 | 1.00 17.37 | A_13 |
|--------------|--------------|------------|------------|--------------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 1388 | N | PRO | 152 | 87.777 | 35.927 | 22.306 | 1.00 10.37 | A_13 |
| ATOM ATOM | 1389 1390 | CD CA | PRO PRO | 152 152 | 86.425 88.030 | 36.450 | 22.575 | 1.00 15.35 | A_13 |
| ATOM | 1391 | CB | PRO | 152 | 86.658 | 34.712 34.412 | 23.087 23.702 | 1.00 11.49 1.00 15.98 | A_13 A_13 |
| ATOM | 1392 | CG | PRO | 152 | 86.083 | 35.789 | 23.898 | 1.00 27.60 | A_13 |
| MOTA | 1393 | C | PRO | 152 | 88.533 | 33.553 | 22.230 | 1.00 18.06 | A_13 |
| ATOM | 1394 | 0 | PRO | 152 | 88.160 | 33.430 | 21.063 | 1.00 16.21 | A_13 |
| MOTA | 1395 | N | ASP | 153 | 89.350 | 32.696 | 22.836 | 1.00 15.86 | A_13 |
| MOTA | 1397 1398 | CA CB | ASP ASP | 153 153 | 89.933 | 31.526 | 22.185 | 1.00 20.25 | A_13 |
| ATOM ATOM | 1399 | CG | ASP | 153 | 90.632 91.843 | 30.630 31.301 | 23.227 23.908 | 1.00 18.17 1.00 24.01 | A_13 A_13 |
| ATOM | 1400 | OD1 | ASP | 153 | 92.517 | 32.159 | 23.284 | 1.00 14.96 | A_13 |
| ATOM | 1401 | OD2 | ASP | 153 | 92.131 | 30.937 | 25.077 | 1.00 20.20 | A_13 |
| MOTA | 1402 | C | ASP | 153 | 88.887 | 30.678 | 21.452 | 1.00 24.64 | A_13 |
| MOTA | 1403 | 0 | ASP | 153 | 89.113 | 30.221 | 20.330 | 1.00 13.51 | A_13 |
| MOTA MOTA | 1404 1406 | N CA | ASP ASP | 154 154 | 87.757 86.664 | 30.453 29.657 | 22.114 21.577 | 1.00 24.11 1.00 19.19 | A_13 |
| ATOM | 1407 | CB | ASP | 154 | 85.527 | 29.632 | 22.587 | 1.00 19.19 | A_13 A_13 |
| ATOM | 1408 | ÇG | ASP | 154 | 84.406 | 28.751 | 22.161 | 1.00 24.26 | A_13 |
| ATOM | 1409 | OD1 | ASP | 154 | 83.314 | 29.291 | 21.950 | 1.00 20.97 | A_13 |
| ATOM | 1410 | OD2 | ASP | 154 | 84.609 | 27.530 | 22.031 | 1.00 20.32 | A_13 |
| MOTA | 1411 | C | ASP | 154 | 86.162 | 30.170 | 20.229 | 1.00 18.99 | A_13 |
| MOTA MOTA | 1412 1413 | O N | ASP ASP | 154 155 | 86.043 85.873 | 29.408 31.465 | 19.277 20.158 | 1.00 22.56 1.00 16.11 | A_13 A_13 |
| ATOM | 1415 | CA | ASP | 155 | 85.407 | 32.078 | 18.917 | 1.00 25.30 | A_13 |
| ATOM | 1416 | CB | ASP | 155 | 85.011 | 33.527 | 19.158 | 1.00 13.32 | A_13 |
| ATOM | 1417 | CG | ASP | 155 | 83.975 | 33.655 | 20.249 | 1.00 11.19 | A_13 |
| MOTA | 1418 | OD1 | ASP | 155 | 84.347 | 34.136 | 21.332 | 1.00 12.26 | A_13 |
| MOTA MOTA | 1419 1420 | OD2 C | ASP ASP | 155 155 | 82.810 86.461 | 33.255 | 20.029 17.828 | 1.00 10.00 1.00 13.98 | A_13 |
| ATOM | 1421 | 0 | ASP | 155 | 86.141 | 31.992 31.656 | 16.687 | 1.00 13.98 | A_13 A_13 |
| ATOM | 1422 | N | VAL | 156 | 87.713 | 32.310 | 18.160 | 1.00 16.49 | A_13 |
| ATOM | 1424 | CA | VAL | 156 | 88.771 | 32.201 | 17.159 | 1.00 27.34 | A_13 |
| MOTA | 1425 | CB | VAL | 156 | 90.145 | 32.826 | 17.625 | 1.00 23.59 | A_13 |
| ATOM | 1426 | CG1 | | 156 | 90.327 | 32.750 | 19.119 | 1.00 13.94 | A_13 |
| MOTA | 1427 1428 | CG2 | | 156 156 | 91.312 | 32.153 | 16.919 | 1.00 21.70 | A_13 |
| MOTA MOTA | 1428 | C | VAL VAL | 156 156 | 88.874 88.946 | 30.738 30.506 | 16.657 15.448 | 1.00 16.95 1.00 13.79 | A_13 A_13 |
| ATOM | 1430 | N | GLN | 157 | 88.762 | 29.763 | 17.561 | 1.00 19.45 | A_13 |
| MOTA | 1432 | CA | GLN | 157 | 88.796 | 28.352 | 17.154 | 1.00 30.53 | A_13 |
| ATOM | 1433 | CB | GLN | 157 | 88.579 | 27.422 | 18.353 | 1.00 23.08 | A'_13 |
| MOTA | 1434 | ÇG | GLN | 157 | 89.633 | 27.521 | 19.452 | 1.00 24.83 | A_13 |
| MOTA MOTA | 1435 1436 | CD OE1 | GLN GLN | 157 157 | 90.950 91.743 | 26.872 27.422 | 19.089 18.316 | 1.00 20.26 1.00 25.80 | A_13 A_13 |
| ATOM | 1437 | NE2 | | 157 | 91.204 | 25.702 | 19.673 | 1.00 23.80 | A_13 |
| MOTA | 1440 | C | GLN | 157 | 87.667 | 28.136 | 16.148 | 1.00 14.16 | A_13 |
| MOTA | 1441 | 0 | GLN | 157 | 87.869 | 27.541 | 15.096 | 1.00 14.11 | A_13 |
| MOTA | 1442 | N | GLY | 158 | 86.505 | 28.709 | 16.437 | 1.00 19.16 | A_13 |
| MOTA MOTA | 1444 1445 | CA C | GLY GLY | 158 158 | 85.361 85.510 | 28.584 29.144 | 15.551 14.143 | 1.00 12.79 1.00 24.46 | A_13 |
| ATOM | 1446 | Ö | GLY | 158 | 85.181 | 28.449 | 13.177 | 1.00 24.40 | A_13 A_13 |
| ATOM | 1447 | N | ILE | 159 | 85.936 | 30.403 | 13.989 | 1.00 22.41 | A_13 |
| ATOM | 1449 | CA | ILE | 159 | 86.091 | 30.946 | 12.628 | 1.00 31.18 | A_13 |
| ATOM | 1450 | CB | ILE | 159 | 86.300 | 32.508 | | 1.00 23.53 | A_13 |
| MOTA MOTA | 1451 1452 | CG2 CG1 | | 159 159 | 84.991 87.022 | 33.203 33.063 | 12.177 | 1.00 17.28 | A_13 |
| MOTA | 1453 | | ILE | 159 | 88.507 | 32.949 | 13.758 13.707 | 1.00 15.28 1.00 14.71 | A_13 A_13 |
| ATOM | 1454 | C | ILE | 159 | 87.226 | 30.280 | 11.875 | 1.00 10.56 | A_13 |
| ATOM | 1455 | 0 | ILE | 159 | 87.167 | | 10.653 | 1.00 18.79 | A_13 |
| MOTA | 1456 | N | GLN | 160 | 88.287 | | | 1.00 20.71 | A_13 |
| MOTA MOTA | 1458 1459 | CA | GLN | 160 | 89.411 | 29.294 | | 1.00 10.00 | A_13 |
| ATOM | 1460 | CB CG | GLN GLN | 160 160 | 90.640 91.114 | | 12.855 13.182 | 1.00 10.00 1.00 13.93 | A_13 A_13 |
| MOTA | 1461 | CD | GLN | 160 | 92.402 | | | 1.00 25.61 | A_13 |
| ATOM | 1462 | 0E1 | | 160 | 92.814 | | | | A_13 |
| MOTA | 1463 | NE2 | 2 GLN | 160 | 93.042 | 31.915 | | | A_13 |
| MOTA | 1466 | С | GLN | 160 | 89.000 | | | | A_13 |
| ATOM | 1467 | O N | GLN | 160 | 89.458 | | | | A_13 |
| MOTA MOTA | 1468 1470 | N CA | SER SER | 161 . 161 | 88.068 87.610 | | | | A_13 A_13 |
| ATOM | 1471 | CB | SER | | 86.688 | | | | A_13 A_13 |
| MOTA | 1472 | OG | SER | | 85.365 | | | | A_13 |
| ATOM | 1474 | C | SER | 161 | 86.913 | 26.048 | | 1.00 26.18 | A_13 |
| ATOM | 1475 | 0 | SER | • | 86.839 | | | | A_13 |
| MOTA MOTA | 1476 1478 | N CA | LEU LEU | | 86.428 85.749 | | | | A_13 |
| 111 OF | 74/0 | CM | ואמת | 102 | 03.747 | 41.433 | 0.000 | 1.00 11.21 | A_13 |

| ATOM 1486 O LEU 162 86.596 27.680 6.584 1.00 15.25 A.1 ATOM 1487 N TWR 163 87.459 29.063 8.135 1.00 26.64 A.1 ATOM 1488 CB TYR 163 88.320 29.796 7.204 1.00 18.28 A.1 ATOM 1488 CB TYR 163 88.320 29.796 7.204 1.00 18.28 A.1 ATOM 1489 CG TYR 163 86.519 31.600 7.039 1.00 18.180 A.1 ATOM 1499 CD LI TYR 163 84.607 31.744 5.754 1.00 10.00 A.1 ATOM 1492 CD LI TYR 163 84.607 31.974 5.503 1.00 16.58 A.1 ATOM 1493 CD TYR 163 84.607 31.974 5.503 1.00 12.00 A.1 ATOM 1494 CZ TYR 163 84.607 31.991 6.576 1.00 11.07 A.1 ATOM 1495 CH TYR 163 88.818 29.669 7.397 1.00 15.63 A.1 ATOM 1497 C TYR 163 88.818 29.669 7.397 1.00 15.67 A.1 ATOM 1498 O TYR 163 88.818 29.669 7.397 1.00 15.67 A.1 ATOM 1498 O TYR 163 89.818 29.669 7.397 1.00 15.67 A.1 ATOM 1498 O TYR 163 89.818 29.669 8.525 1.00 18.34 A.1 ATOM 1502 C GLY 164 90.225 29.096 8.526 1.00 18.34 A.1 ATOM 1503 O GLY 164 90.225 29.096 8.526 1.00 16.6 A.1 ATOM 1503 O GLY 164 91.334 31.139 9.757 1.00 21.42 A.1 ATOM 1504 OT GLY 164 93.153 30.250 9.858 1.00 21.99 A.1 ATOM 3010 ON EN 166 93.153 30.250 9.858 1.00 21.99 A.1 ATOM 3010 ON EN 166 65.514 41.122 10.661 1.00 15.63 A.1 ATOM 3010 CR LI 164 93.153 30.250 9.858 1.00 21.99 A.1 ATOM 3010 CR LI 164 93.153 30.250 9.858 1.00 21.99 A.1 ATOM 3010 CR LI 164 93.153 30.250 9.858 1.00 21.99 A.1 ATOM 3010 CR LI 169 66.626 35.60 1.00 16.6 A.1 ATOM 3010 CR LI 169 66.626 35.60 1.00 16.60 A.1 ATOM 3010 CR LI 169 66.626 35.60 1.00 16.60 A.1 ATOM 3010 CR LI 169 67.400 35.999 20.267 1.00 38.86 A.66 ATOM 3021 CR LI 169 67.400 35.999 20.267 1.00 38.86 A.66 ATOM 3021 CR LI 169 67.400 35.999 20.267 1.00 38.86 A.67 ATOM 3022 CR WAY 169 68.866 34.739 22.265 1.00 18.83 A.67 ATOM 3022 CR WAY 169 68.866 34.739 22.265 1.00 18.83 A.67 ATOM 3022 CR WAY 169 69.699 36.617 21.141 1.00 33.16 A.66 ATOM 3022 CR WAY 169 69.999 36.617 21.141 1.00 33.16 A.66 ATOM 3022 CR WAY 169 66.866 36.75 22.21 21.80 1.00 32.16 A.67 ATOM 3022 CR WAY 169 66.866 36.75 22.21 21.00 22.66 B.67 ATOM 3022 CR WAY 169 66.866 34.739 22.265 1.00 18.82 ATOM 3030 CR WAY 169 66.866 34.739 22.265 1.00 | ATOM ATOM ATOM AOTA | 1479 1480 1481 1482 | CB LEU CG LEU CD1 LEU CD2 LEU | 162 162 162 162 | 84.584 83.489 82.596 82.672 | 28.477 28.144 29.351 26.949 | 9.007 10.021 10.217 9.548 | 1.00 14.37 1.00 31.09 1.00 14.96 1.00 23.87 | A_13 A_13 A_13 A_13 |
|--|------------------------------|------------------------------|--|--------------------------|--------------------------------------|--------------------------------------|------------------------------------|--|------------------------------|
| ATOM 1485 N TYR 163 87.459 29.063 8.135 1.00 26.64 AL ATOM 1487 CA TYR 163 88.320 29.796 7.207 1.00 26.89 AL ATOM 1488 CB TYR 163 87.977 31.289 7.277 1.00 26.89 AL ATOM 1499 CG TYR 163 86.519 31.600 7.039 1.00 18.80 AL ATOM 1491 CEL TYR 163 86.519 31.600 7.039 1.00 18.80 AL ATOM 1491 CEL TYR 163 86.527 31.744 5.749 1.00 10.03 AL ATOM 1491 CEL TYR 163 86.527 31.744 5.749 1.00 10.03 AL ATOM 1492 CDZ TYR 163 86.527 31.67Z 8.099 1.00 16.58 AL ATOM 1493 CEZ TYR 163 86.526 31.867 7.873 1.00 12.23 AL ATOM 1493 CEZ TYR 163 84.266 31.867 7.873 1.00 12.23 AL ATOM 1495 CD TYR 163 88.522 31.67Z 8.099 1.00 16.58 AL ATOM 1495 CD TYR 163 88.5622 31.67Z 8.099 1.00 16.576 AL ATOM 1498 CD TYR 163 80.590 20.685 7.357 1.00 12.793 AL ATOM 1498 CD TYR 163 80.590 20.685 7.357 1.00 12.793 AL ATOM 1498 CD TYR 163 80.590 20.685 7.357 1.00 12.793 AL ATOM 1498 CD TYR 163 80.590 20.685 8.226 1.00 18.34 AL ATOM 1501 CA GLY 164 91.636 28.966 8.226 1.00 18.34 AL ATOM 1502 CD GLY 164 91.636 28.966 8.226 1.00 18.34 AL ATOM 1503 CD GLY 164 91.334 31.139 9.775 1.00 21.42 AL ATOM 1503 CD GLY 164 91.334 31.139 9.775 1.00 21.42 AL ATOM 1503 CD GLY 164 91.334 31.139 9.775 1.00 21.42 AL ATOM 3009 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATOM 3009 ZN ZN 166 673.275 35.223 18.371 1.00 27.40 ATOM 3010 ZN ZN 167 65.511 41.122 10.564 1.00 27.40 ATOM 3012 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 66.26 35.606 31.607 1.00 38.86 A65 ATOM 3012 CA CA 168 66.26 35.606 31.607 1.00 38.86 A65 ATOM 3012 CA CA 168 66.26 35.606 31.007 1.00 31.72 AC ATOM 3012 CA CA 168 66.26 35.606 31.007 1.00 31.72 AC ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35.606 31.00 31.70 ATOM 3012 CA CA 168 66.26 35 | | | | | | | | | A_13 A_13 |
| ATOM 1488 CB TYR 163 87.977 31.289 7.277 1.00 26.89 A_I ATOM 1499 CG TYR 163 86.519 31.600 7.091 1.00 18.80 A_I ATOM 1491 CEL TYR 163 86.527 31.744 5.749 1.00 10.00 10.01 A_I ATOM 1491 CEL TYR 163 86.027 31.744 5.749 1.00 10.02 83 A_I ATOM 1492 CD2 TYR 163 85.622 31.672 8.099 1.00 16.58 A_I ATOM 1493 CEZ TYR 163 88.262 31.672 8.099 1.00 16.58 A_I ATOM 1493 CEZ TYR 163 88.262 31.867 7.873 1.00 12.32 A_I ATOM 1495 CH TYR 163 82.472 32.141 6.331 1.00 21.93 A_I ATOM 1495 CH TYR 163 82.472 32.141 6.331 1.00 21.93 A_I ATOM 1497 C TYR 163 89.518 29.669 7.391 1.00 15.57 A_I ATOM 1497 C TYR 163 89.518 29.669 7.391 1.00 15.57 A_I ATOM 1499 N CTYR 163 89.518 29.669 7.392 1.00 15.67 A_I ATOM 1499 N CTYR 163 89.518 29.669 6.826 1.00 18.42 A_I ATOM 1501 CA GLY 164 99.536 29.968 6.826 1.00 18.42 A_I ATOM 1501 CA GLY 164 99.536 29.968 6.826 1.00 18.42 A_I ATOM 1502 C GLY 164 99.534 31.139 9.775 1.00 21.42 A_I ATOM 1503 O GLY 164 99.3353 30.250 9.858 1.00 21.42 A_I ATOM 3009 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATOM 3009 ZN ZN 166 673.275 35.223 18.371 1.00 27.40 ATOM 3012 CA CA 168 64.855 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 64.855 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 168 66.263 50.60 19.140 1.00 1.78 ATOM 3012 CA CA 168 66.263 50.60 19.140 1.00 1.78 ATOM 3012 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 66.26 35.60 19.161 1.00 30.96 A6 ATOM 3021 CA CA 168 67.90 19.10 | MOTA | 1485 | N TYR | 163 | 87.459 | 29.063 | 8.135 | 1.00 26.64 | A_13 |
| ATOM 1489 CG TYR 163 86.519 31.600 7.039 1.00 18.80 A_1 ATOM 1491 CBL TYR 163 86.027 31.744 5.749 1.00 10.00 ATOM 1491 CBL TYR 163 84.680 31.936 5.515 1.00 12.83 A_1 ATOM 1492 CD2 TYR 163 84.266 31.867 7.873 1.00 12.83 A_1 ATOM 1493 CZ TYR 163 84.266 31.867 7.873 1.00 12.32 A_1 ATOM 1495 CD4 TYR 163 82.472 32.141 6.331 1.00 12.32 A_1 ATOM 1495 CD4 TYR 163 82.472 32.141 6.331 1.00 12.32 A_1 ATOM 1495 CD4 TYR 163 82.472 32.141 6.331 1.00 12.32 A_1 ATOM 1495 CD4 TYR 163 89.818 29.669 7.397 1.00 12.52 A_1 ATOM 1498 N GLY 164 90.225 29.096 8.256 1.00 18.92 A_1 ATOM 1499 N GLY 164 90.225 29.096 8.256 1.00 18.94 A_1 ATOM 1501 C GLY 164 91.636 28.9696 8.826 1.00 10.61 A_1 ATOM 1501 C GLY 164 91.636 28.9697 3.95 1.00 10.61 A_1 ATOM 1501 C GLY 164 93.333 31.931 0.50 9.95 1.00 10.61 A_1 ATOM 3010 C GLY 164 93.333 31.050 2.97 9.95 8.85 1.00 21.92 A_1 ATOM 3010 C GLY 164 93.333 31.050 2.97 9.858 1.00 22.9 A_2 ATOM 3010 C GLY 164 93.333 31.050 2.97 9.858 1.00 22.9 A_2 ATOM 3010 C GLY 164 93.333 31.050 2.97 9.858 1.00 22.9 A_2 ATOM 3010 C GLY 164 93.333 31.93 9.377 1.854 1.00 21.92 A_2 ATOM 3010 C GLY 164 93.333 9.377 1.854 1.00 21.92 A_2 ATOM 3010 C GLY 164 93.333 9.377 1.854 1.00 40.73 ATOM 3010 C GLY 164 93.335 99.377 1.854 1.00 40.73 ATOM 3010 C GLY 164 93.856 91.30 0.30 9.9 8.85 1.00 22.9 A_2 ATOM 3010 C GLY 164 93.333 99.377 1.854 1.00 40.73 ATOM 3010 C C C WAY 169 66.263 35.606 19.161 1.00 30.96 A65 ATOM 3020 C C WAY 169 66.263 35.606 19.161 1.00 30.96 A65 ATOM 3020 C C WAY 169 66.856 35.606 19.161 1.00 30.96 A65 ATOM 3020 C C WAY 169 68.861 35.602 31.10 31.1 1.00 21.78 A65 ATOM 3021 C C WAY 169 68.861 35.602 31.10 31.1 1.00 21.78 A65 ATOM 3022 C WAY 169 68.861 35.602 31.72 A60 31.1 1.00 31.73 A66 ATOM 3022 C WAY 169 68.861 35.602 31.700 30.1 1.00 31.73 A66 ATOM 3024 CD WAY 169 68.861 35.602 31.700 30.1 1.00 31.71 A66 ATOM 3026 C WAY 169 68.861 35.602 31.700 30.1 1.00 31.71 A66 ATOM 3026 C WAY 169 68.861 35.602 31.700 30.1 1.00 31.72 A66 ATOM 3026 C WAY 169 68.861 35.500 20.07 1.00 30.96 A66 ATOM 3026 C WAY 169 68.861 | | | | | | | | | A_13 A_13 |
| ATOM 1491 CEL TYR 163 84.680 31.936 5.515 1.00 12.83 A_L ATOM 1493 CEZ TYR 163 85.622 31.672 8.099 1.00 16.58 ATOM 1494 CZ TYR 163 84.266 31.867 7.873 1.00 12.32 A_L ATOM 1495 OH TYR 163 82.472 32.141 6.331 1.00 21.93 A_L ATOM 1495 OH TYR 163 82.472 32.141 6.331 1.00 21.93 A_L ATOM 1495 OH TYR 163 89.818 29.669 7.397 1.00 15.67 A_L ATOM 1498 O TYR 163 90.590 30.089 6.526 1.00 18.92 A_L ATOM 1499 N GLY 164 91.636 28.966 8.825 1.00 18.94 A_L ATOM 1501 CA GLY 164 91.636 28.966 8.826 1.00 10.61 A_L ATOM 1502 C GLY 164 91.636 28.966 8.826 1.00 10.61 A_L ATOM 1504 OT GLY 164 91.334 31.139 9.775 1.00 21.42 A_L ATOM 1504 OT GLY 164 91.334 31.139 9.775 1.00 21.42 A_L ATOM 3009 2N ZN 166 73.275 35.223 18.371 1.00 27.40 ATC ATOM 3010 2N ZN 166 73.275 35.223 18.371 1.00 27.40 ATC ATOM 3011 CA CA 168 64.285 44.152 21.654 1.00 17.66 ATC ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATC ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATC ATOM 3018 CF1 WAY 169 67.400 35.999 20.267 1.00 38.86 AGS ATOM 3018 CF1 WAY 169 67.400 35.999 20.267 1.00 38.86 AGS ATOM 3021 CA CA 169 67.999 35.400 17.901 1.00 41.17 AGS ATOM 3021 CA CA 169 67.999 35.400 17.901 1.00 41.17 AGS ATOM 3022 C4 WAY 169 68.961 55.601 19.161 1.00 30.96 AGS ATOM 3022 C4 WAY 169 68.961 55.623 17.728 1.00 33.71 AGS ATOM 3022 C4 WAY 169 68.961 63.267 22.189 1.00 22.78 AGS ATOM 3022 C4 WAY 169 68.966 34.739 22.685 1.00 25.69 AGS ATOM 3023 NZO WAY 169 68.966 34.739 22.685 1.00 25.69 AGS ATOM 3026 C2 WAY 169 68.967 33.93 38.039 18.811 1.00 35.73 AGS ATOM 3026 C2 WAY 169 68.967 39.93 37.71 AGS AGS ATOM 3027 C7 WAY 169 69.939 36.617 21.141 1.00 33.16 AGS ATOM 3028 CZ WAY 169 69.939 36.617 21.141 1.00 33.16 AGS ATOM 3028 CZ WAY 169 69.939 36.617 21.141 1.00 33.16 AGS ATOM 3028 CZ WAY 169 69.939 36.617 21.141 1.00 33.16 AGS ATOM 3028 CZ WAY 169 69.939 37.70 AGS AGS AGS ATOM 3028 CZ WAY 169 69.939 37.70 AGS | MOTA | | | | | | | | A_13 |
| ATOM 1493 CEZ TYR 163 84.266 31.867 7.873 1.00 12.32 A_1 ATOM 1495 OH TYR 163 83.807 31.991 6.576 1.00 11.77 ATOM 1495 OH TYR 163 82.472 32.141 6.331 1.00 21.93 A_1 ATOM 1495 OH TYR 163 89.818 29.669 7.397 1.00 15.67 ATOM 1498 O TYR 163 90.590 30.089 6.526 1.00 18.92 A_1 ATOM 1499 N GUY 164 90.225 29.096 8.526 1.00 18.94 A_1 ATOM 1501 CA GUY 164 91.636 28.966 8.826 1.00 10.61 A_1 ATOM 1502 C GUY 164 91.636 28.966 8.826 1.00 10.61 A_1 ATOM 1503 O GUY 164 91.334 31.139 9.775 1.00 21.42 A_1 ATOM 1503 O GUY 164 91.334 31.139 9.775 1.00 21.42 A_1 ATOM 3009 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3011 CA CA 168 64.285 44.152 21.655 1.00 11.76 ATO ATOM 3011 CA CA 168 64.285 44.152 21.655 1.00 11.76 ATO ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATO ATOM 3018 CFI WAY 169 67.400 35.999 20.267 1.00 38.86 A65 ATOM 3019 CH WAY 169 67.400 35.999 20.267 1.00 38.86 A65 ATOM 3020 CZ WAY 169 68.561 35.601 19.161 1.00 30.96 A65 ATOM 3021 C3 WAY 169 67.999 35.600 17.901 1.00 41.17 A65 ATOM 3022 C4 WAY 169 68.857 135.623 17.728 1.00 33.71 A66 ATOM 3022 C4 WAY 169 68.867 36.216 20.078 1.00 33.71 A66 ATOM 3022 C4 WAY 169 68.867 36.216 20.078 1.00 33.71 A66 ATOM 3022 C4 WAY 169 68.867 36.216 20.078 1.00 33.71 A66 ATOM 3022 C4 WAY 169 68.867 35.621 20.078 1.00 33.71 A66 ATOM 3023 WAO WAY 169 68.867 35.621 20.078 1.00 33.71 A66 ATOM 3023 CC 20 WAY 169 68.867 35.621 20.078 1.00 33.71 A66 ATOM 3025 C23 WAY 169 68.867 35.622 17.728 1.00 33.71 A66 ATOM 3027 C7 WAY 169 68.867 35.622 17.758 1.00 20.288 A66 ATOM 3027 C7 WAY 169 68.867 36.216 20.078 1.00 33.71 A66 ATOM 3028 CB WAY 169 69.339 38.639 18.811 1.00 35.73 A66 ATOM 3027 C7 WAY 169 69.339 36.639 18.811 1.00 35.73 A66 ATOM 3028 CB WAY 169 69.399 36.617 21.141 1.00 33.16 A66 ATOM 3029 CC WAY 169 69.399 36.617 21.141 1.00 33.16 A66 ATOM 3029 CC WAY 169 69.399 36.617 21.141 1.00 33.16 A66 ATOM 3029 CC WAY 169 69.399 36.617 21.141 1.00 33.16 A66 ATOM 3029 CC WAY 169 69.6896 34.739 22.2685 1.00 20.288 A66 ATOM 3029 C | MOTA | 1491 | CE1 TYR | 163 | | 31.936 | 5.515 | 1.00 12.83 | A_13 |
| ATOM 1494 CZ TYR 163 83.807 31.991 6.576 1.00 11.77 A_I ATOM 1497 C TYR 163 82.472 32.141 6.331 1.00 21.93 A_I ATOM 1497 C TYR 163 89.818 29.669 7.397 1.00 15.67 A_I ATOM 1498 O TYR 163 89.818 29.669 7.397 1.00 15.67 A_I ATOM 1499 N GLY 164 90.225 29.096 8.525 1.00 18.92 A_I ATOM 1501 CA GLY 164 91.636 28.966 8.966 8.262 1.00 18.92 A_I ATOM 1502 C GLY 164 91.636 28.966 8.966 N.625 1.00 18.92 A_I ATOM 1503 O GLY 164 92.149 30.215 9.525 1.00 15.63 A_I ATOM 1503 O GLY 164 91.334 11.39 9.775 1.00 21.42 A_I ATOM 1504 OT GLY 164 93.353 30.250 9.858 1.00 21.99 A_I ATOM 300 0 ZN ZN 166 73.275 55.223 18.371 1.00 27.96 A_I ATOM 3010 ZN ZN 167 65.511 41.122 10.564 1.00 27.96 A_I ATOM 3010 ZN ZN 167 65.511 41.122 10.564 1.00 27.96 A_I ATOM 3011 CA CA 168 64.285 44.152 21.655 1.00 11.76 A_I ATOM 3012 CA CA 165 64.285 44.152 21.655 1.00 11.76 A_I ATOM 3012 CA CA 165 67.301 93.93.77 1.854 1.00 40.73 A_I ATOM 3013 CT WAY 169 66.626 35.606 19.161 1.00 30.96 A_I ATOM 3019 CH WAY 169 66.626 35.606 19.161 1.00 30.96 A_I ATOM 3012 CA WAY 169 66.526 35.606 19.161 1.00 30.96 A_I ATOM 3012 CA WAY 169 68.807 36.039 18.811 1.00 31.73 A_I ATOM 3012 CX WAY 169 68.807 36.039 18.811 1.00 31.73 A_I ATOM 3012 CX WAY 169 68.807 36.216 20.078 1.00 33.71 A_I ATOM 3012 CX WAY 169 68.807 36.216 20.078 1.00 33.71 A_I ATOM 3012 CX WAY 169 68.807 36.216 20.078 1.00 33.71 A_I ATOM 3012 CX WAY 169 68.939 36.617 21.141 1.00 30.96 A_I ATOM 3012 CX WAY 169 68.969 3.39 36.617 21.141 1.00 30.95 A_I ATOM 3012 CX WAY 169 68.960 39 36.617 21.141 1.00 30.95 A_I ATOM 3012 CX WAY 169 68.960 39 36.617 21.141 1.00 30.95 A_I ATOM 3012 CX WAY 169 68.960 39 36.617 21.141 1.00 31.73 A_I ATOM 3013 CX WAY 169 68.960 39 30.32 C | | | | | | | | | A_13 A_13 |
| ATOM 1498 O TYR 163 89.818 29.669 7.397 1.00 15.67 A_I ATOM 1498 O TYR 163 90.590 30.089 6.526 1.00 18.92 A_I ATOM 1499 N GLY 164 90.225 29.996 8.525 1.00 18.92 A_I ATOM 1501 CA GLY 164 91.636 28.966 8.525 1.00 10.61 A_I ATOM 1502 C GLY 164 91.636 28.966 8.826 1.00 10.61 A_I ATOM 1503 O GLY 164 91.334 31.139 9.775 1.00 15.63 A_I ATOM 1504 OT GLY 164 91.334 31.139 9.755 1.00 21.42 A_I ATOM 309 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 A_I ATOM 3010 ZN ZN 166 65.511 41.122 10.564 1.00 27.86 A_I ATOM 3011 CA CA 168 64.285 44.152 21.635 1.00 41.76 A_I ATOM 3012 CA CA 168 64.285 44.152 21.635 1.00 40.73 A_I ATOM 3012 CA CA 168 73.319 39.377 1.854 1.00 40.73 A_I ATOM 3018 CF1 WAY 169 66.626 35.606 19.161 1.00 30.96 A_I ATOM 3019 CH WAY 169 67.199 35.400 17.901 1.00 41.17 A_I ATOM 3020 C2 WAY 169 68.361 35.623 17.728 1.00 36.26 A_I ATOM 3021 C3 WAY 169 68.851 35.623 17.928 1.00 35.73 A_I ATOM 3021 C3 WAY 169 68.867 36.216 20.078 1.00 33.71 A_I ATOM 3022 C4 WAY 169 68.867 36.216 20.078 1.00 33.71 A_I ATOM 3023 N20 WAY 169 68.869 36.617 21.141 1.00 35.73 A_I ATOM 3025 C22 WAY 169 68.969 36.617 21.141 1.00 33.16 A_I ATOM 3026 C28 WAY 169 68.869 36.617 21.141 1.00 31.72 A_I ATOM 3027 C27 WAY 169 68.969 34.379 22.685 1.00 25.69 A_I ATOM 3028 C20 WAY 169 68.969 34.739 22.685 1.00 25.69 A_I ATOM 3020 C24 WAY 169 68.969 37.39 32.748 22.265 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 67.743 32.2685 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 77.733 32.748 22.265 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 22.265 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 22.265 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 22.265 1.00 25.69 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 22.265 1.00 25.96 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 22.265 1.00 25.96 A_I ATOM 3030 C24 WAY 169 69.699 35.617 21.414 1.00 16.09 A_I ATOM 3030 C24 WAY 169 69.699 35.617 21.414 1.00 16.99 A_I ATOM 3030 C24 WAY 169 68.969 77.73 32.274 1.00 22.94 A_I ATOM 3030 C24 WAY 169 68.899 78.617 32.286 1.00 22.96 A_I ATOM 3030 | MOTA | 1494 | CZ TYR | 163 | 83.807 | 31.991 | 6.576 | 1.00 11.77 | A_13 |
| ATOM 1498 O TYR 163 90.590 30.089 6.526 1.00 18.92 A_1 ATOM 1499 N GLY 164 91.636 28.966 8.525 1.00 18.34 A_1 ATOM 1501 CA GLY 164 91.636 28.966 8.525 1.00 18.34 A_1 ATOM 1502 C GLY 164 92.149 30.215 9.525 1.00 15.63 A_1 ATOM 1503 O GLY 164 92.149 30.215 9.525 1.00 15.63 A_1 ATOM 1503 O GLY 164 93.353 30.250 9.858 1.00 21.99 A_1 ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.86 ATO ATOM 3011 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATO ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATO ATOM 3018 CF1 WAY 169 66.626 35.606 19.161 1.00 30.96 A66 ATOM 3018 CF1 WAY 169 66.626 35.606 19.161 1.00 30.96 A66 ATOM 3019 CH WAY 169 67.199 35.400 17.901 1.00 41.17 A66 ATOM 3020 CZ WAY 169 68.851 35.623 17.728 1.00 36.26 A65 ATOM 3021 C3 WAY 169 68.850 36.019 18.811 1.00 35.73 A66 ATOM 3022 C4 WAY 169 68.850 36.216 20.078 1.00 33.16 A66 ATOM 3022 C4 WAY 169 68.850 36.216 20.078 1.00 33.16 A66 ATOM 3023 N20 WAY 169 68.850 36.212 21.141 1.00 33.16 A66 ATOM 3025 C23 WAY 169 68.850 36.212 21.141 1.00 33.16 A66 ATOM 3026 C28 WAY 169 68.850 34.739 22.685 1.00 25.69 A66 ATOM 3027 C27 WAY 169 68.850 37.35 A60 ATOM 3028 CM WAY 169 68.980 36.212 21.89 1.00 29.78 A66 ATOM 3028 CM WAY 169 68.980 38.617 221.141 1.00 33.16 A66 ATOM 3026 C28 WAY 169 68.980 38.617 221.141 1.00 33.16 A66 ATOM 3030 C24 WAY 169 68.980 38.617 221.141 1.00 33.16 A66 ATOM 3030 C24 WAY 169 68.980 38.617 221.141 1.00 33.61 A66 ATOM 3030 C24 WAY 169 68.980 38.617 221.141 1.00 33.61 A66 ATOM 3031 S21 WAY 169 67.141 34.238 24.205 1.00 31.72 A66 ATOM 3031 S21 WAY 169 68.990 36.617 221.141 1.00 33.61 A66 ATOM 3030 C24 WAY 169 70.37 35.640 22.189 1.00 29.78 A66 ATOM 3031 S21 WAY 169 68.900 38.617 221.141 1.00 33.61 A66 ATOM 3031 S21 WAY 169 69.900 38.617 221.141 1.00 38.61 A66 ATOM 3031 C24 WAY 169 68.800 38.8 | | | | | | | | | A_13 A_13 |
| ATOM 1501 CA GLY 164 91.636 28.966 8.826 1.00 10.61 A.I ATOM 1502 C GLY 164 92.149 30.215 9.525 1.00 15.63 A.I ATOM 1503 O GLY 164 93.134 31.139 9.775 1.00 21.42 A.I ATOM 1504 OT GLY 164 93.153 30.250 9.858 1.00 21.99 A.I ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATC ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATC ATOM 3010 ZN ZN 166 75.551 41.122 10.564 1.00 27.86 ATC ATOM 3011 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATC ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 11.76 ATC ATOM 3012 CR CA 165 73.319 39.377 1.854 1.00 11.76 ATC ATOM 3017 C5 WAY 169 67.400 35.999 20.267 1.00 38.86 A68 ATOM 3018 CF1 WAY 169 67.400 35.999 20.267 1.00 38.86 A68 ATOM 3019 CH WAY 169 67.400 35.999 20.267 1.00 30.96 A66 ATOM 3020 C2 WAY 169 68.551 35.623 17.728 1.00 35.73 A66 ATOM 3021 C3 WAY 169 68.551 35.623 17.728 1.00 33.71 A66 ATOM 3022 C4 WAY 169 68.807 36.216 20.078 1.00 33.71 A66 ATOM 3023 N20 WAY 169 68.807 36.517 21.141 1.00 33.16 A66 ATOM 3020 C2 WAY 169 68.807 36.517 21.141 1.00 33.16 A66 ATOM 3020 C2 WAY 169 68.807 36.517 21.141 1.00 33.16 A66 ATOM 3020 C2 WAY 169 68.986 34.739 22.685 1.00 25.69 A66 ATOM 3020 C22 WAY 169 68.986 34.739 22.685 1.00 25.69 A66 ATOM 3020 C22 WAY 169 68.986 34.739 22.685 1.00 25.69 A66 ATOM 3020 C22 WAY 169 67.743 32.328 22.266 1.00 27.88 A67 ATOM 3020 C24 WAY 169 66.975 33.213 21.577 1.00 24.43 A67 ATOM 3030 C24 WAY 169 67.743 32.328 22.266 1.00 27.88 A67 ATOM 3030 C24 WAY 169 69.797 33.2748 22.266 1.00 22.96 A67 ATOM 3030 C24 WAY 169 69.797 38.233 21.577 1.00 24.43 A67 ATOM 3030 C24 WAY 169 69.797 38.233 21.577 1.00 24.43 A67 ATOM 3031 C31 WAY 169 77.738 38.564 22.016 1.00 27.88 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.216 1.00 11.82 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.206 1.00 23.96 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.206 1.00 23.29 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.206 1.00 23.29 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.206 1.00 23.29 A67 ATOM 3030 C31 WAY 169 77.738 38.564 22.206 1.00 23.29 A67 ATOM 3030 C31 WAY 169 78.789 A67 ATOM 3030 C31 | | | | | | | | | A_13 |
| ATOM 1503 O GLY 164 91.334 31.139 9.775 1.00 21.42 A.1 ATOM 1504 OT GLY 164 91.335 30.250 9.858 1.00 21.99 A.1 ATOM 3009 2N ZN 166 73.275 35.223 18.371 1.00 27.40 ATO 3010 ZN ZN 167 65.511 41.122 10.564 1.00 27.86 ATO ATOM 3011 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 ATOM ATOM 3012 CR CA 165 73.319 39.377 1.854 1.00 40.73 ATOM ATOM 3012 CR CA 165 73.319 39.377 1.854 1.00 40.73 ATOM ATOM 3018 CF1 WAY 169 66.263 35.606 19.161 1.00 30.96 AGO ATOM 3019 CH WAY 169 66.263 35.606 19.161 1.00 30.96 AGO ATOM 3019 CH WAY 169 66.562 35.506 19.161 1.00 30.96 AGO ATOM 3021 C3 WAY 169 68.561 35.623 17.728 1.00 36.26 AGO ATOM 3021 C3 WAY 169 69.339 36.039 18.811 1.00 35.73 AGO ATOM 3022 C4 WAY 169 69.399 36.216 20.078 1.00 33.71 AGO ATOM 3023 N20 WAY 169 69.899 36.617 21.141 1.00 33.16 AGO ATOM 3023 N20 WAY 169 69.899 36.617 21.141 1.00 33.16 AGO ATOM 3025 C23 WAY 169 68.807 35.640 22.189 1.00 29.78 AGO ATOM 3025 C23 WAY 169 68.807 35.640 22.189 1.00 29.78 AGO ATOM 3026 C28 WAY 169 66.986 34.739 22.665 1.00 31.72 AGO ATOM 3027 C27 WAY 169 66.991 33.640 22.189 1.00 29.78 AGO ATOM 3028 CM WAY 169 66.991 33.061 23.490 1.00 32.16 AGO ATOM 3029 N25 WAY 169 67.703 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 67.703 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 67.703 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 67.703 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 67.73 33.546 22.016 1.00 27.88 AGO ATOM 3030 C24 WAY 169 67.73 33.546 22.016 1.00 27.88 AGO ATOM 3030 C24 WAY 169 67.73 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 67.73 32.748 22.425 1.00 42.39 AGO ATOM 3030 C24 WAY 169 69.757 38.213 21.577 1.00 24.43 AGO ATOM 3030 C24 WAY 169 77.523 39.163 20.259 1.00 13.26 AGO ATOM 3030 C24 WAY 169 69.757 38.232 22.257 1.00 23.99 AGO ATOM 3030 C24 WAY 169 77.523 39.453 20.259 1.00 29.75 AGO ATOM 3030 C24 WAY 169 77.523 39.453 20.259 1.00 29.75 AGO ATOM 3030 C24 WAY 169 69.759 38.253 1.00 20.65 AGO ATOM 303 | MOTA | 1501 | CA GLY | 164 | 91.636 | 28.966 | 8.826 | 1.00 10.61 | A_13 |
| ATOM 3009 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3010 ZN ZN 166 73.275 35.223 18.371 1.00 27.40 ATO ATOM 3011 CR CA 168 64.285 44.152 10.564 1.00 27.86 ATO ATOM 3011 CR CA 168 64.285 44.152 21.635 1.00 11.76 ATOM ATOM 3012 CR CA 165 73.319 39.377 1.854 1.00 40.73 ATOM 3013 CR CR WAY 169 67.400 35.99 20.267 1.00 38.86 ASO ATOM 3019 CH WAY 169 66.266 35.606 19.161 1.00 30.96 ASO ATOM 3020 CZ WAY 169 66.266 35.606 19.161 1.00 30.96 ASO ATOM 3020 CZ WAY 169 68.561 35.603 17.728 1.00 36.266 ASO ATOM 3022 C4 WAY 169 68.865 1 35.603 17.728 1.00 36.266 ASO ATOM 3022 C4 WAY 169 68.807 36.216 20.078 11.00 33.16 ASO ATOM 3022 C4 WAY 169 68.807 36.216 20.078 11.00 33.16 ASO ATOM 3024 CD WAY 169 68.807 36.216 20.078 11.00 33.16 ASO ATOM 3024 CD WAY 169 68.806 34.739 22.685 1.00 25.69 ASO ATOM 3025 C23 WAY 169 68.806 34.739 22.685 1.00 25.69 ASO ATOM 3026 C28 WAY 169 68.806 34.739 22.685 1.00 25.69 ASO ATOM 3028 CM WAY 169 66.806 38.309 23.798 1.00 31.72 ASO ATOM 3028 CM WAY 169 66.806 38.309 23.798 1.00 31.72 ASO ATOM 3028 CM WAY 169 66.9139 33.540 22.189 1.00 32.768 ASO ATOM 3028 CM WAY 169 66.929 33.046 22.016 1.00 27.88 ASO ATOM 3030 C24 WAY 169 66.921 33.061 23.490 1.00 32.16 ASO ATOM 3031 S21 WAY 169 66.921 33.061 23.490 1.00 32.16 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 20.269 1.00 32.80 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.757 38.213 21.577 1.00 24.43 ASO ATOM 3033 C21 WAY 169 69.49 39.453 20.169 1.00 11.82 ASO ATOM 3030 C24 WAY 169 69.49 39.453 20.169 1.00 11.82 ASO ATOM 3033 C21 WAY 169 69.49 39.453 20.169 1.00 11.82 ASO ATOM 3034 C20 WAY 169 67.60 ASO ATOM 30.40 C20 WAY 169 69.49 39.453 20.169 1.00 11.88 ASO ATOM 3034 C20 WAY 169 69.60 ASO ATOM 30.90 ASO ATOM 30.90 ASO | | | | | | | | | A_13 A_13 |
| ATOM 3010 2N ZN 167 65.511 41.122 10.564 1.00 27.86 ATOM 3011 CA CA 168 64.285 44.152 21.635 1.00 11.76 ATC ATOM 3012 CA CA 1655 73.319 39.377 1.854 1.00 40.73 ATC ATOM 3017 C5 WAY 169 67.400 35.99 20.267 1.00 38.86 AS ATOM 3018 CF1 WAY 169 66.626 35.606 19.161 1.00 30.96 AG ATOM 3019 CH WAY 169 66.526 35.606 19.161 1.00 30.96 AG ATOM 3020 C2 WAY 169 68.561 35.602 317.728 1.00 36.26 AG ATOM 3021 C3 WAY 169 68.561 35.623 17.728 1.00 36.26 AG ATOM 3022 C4 WAY 169 68.807 36.231 C2.078 11.00 33.16 AG ATOM 3022 C4 WAY 169 68.807 36.232 C2.078 11.00 33.71 AG ATOM 3022 C4 WAY 169 68.807 36.216 20.078 11.00 33.16 AG ATOM 3024 CD WAY 169 68.807 36.617 21.141 1.00 33.16 AG ATOM 3024 CD WAY 169 68.806 34.739 22.665 1.00 25.69 AG ATOM 3025 C23 WAY 169 68.806 34.739 22.665 1.00 25.69 AG ATOM 3026 C28 WAY 169 66.8186 34.739 22.665 1.00 33.61 AG ATOM 3027 C27 WAY 169 67.141 34.238 24.205 1.00 33.61 AG ATOM 3028 CM WAY 169 66.929 33.0617 23.490 1.00 32.16 AG ATOM 3022 NO WAY 169 66.929 33.061 23.490 1.00 32.16 AG ATOM 3022 C4 WAY 169 66.929 33.064 22.189 1.00 25.69 AG ATOM 3033 C21 WAY 169 66.929 33.064 22.089 1.00 31.72 AG ATOM 3033 C21 WAY 169 66.97 70.03 32.748 22.426 1.00 42.39 AG ATOM 3030 C24 WAY 169 66.929 33.546 22.016 1.00 27.88 AG ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 AG ATOM 3033 C21 WAY 169 77.513 38.570 21.438 1.00 29.69 AG ATOM 3033 C21 WAY 169 77.503 39.453 20.169 1.00 11.82 AG ATOM 3033 C21 WAY 169 77.503 39.453 20.169 1.00 11.82 AG ATOM 3033 C31 WAY 169 77.503 39.453 20.169 1.00 11.82 AG ATOM 3033 C31 WAY 169 77.503 39.453 20.169 1.00 11.82 AG ATOM 3030 C24 WAY 169 77.504 39.509 22.271 1.00 12.69 AG ATOM 3030 C4 WAY 169 77.504 39.509 20.607 1.00 26.57 AG ATOM 3030 C4 WAY 169 77.504 39.509 20.607 1.00 26.57 AG ATOM 3030 C4 WAY 169 77.504 39.509 20.271 1.00 11.82 AG ATOM 3030 C4 WAY 169 77.504 39.509 20.271 1.00 11.82 AG ATOM 3030 C4 WAY 169 79.504 39.509 20.271 1.00 12.69 AG ATOM 3030 C4 WAY 169 79.504 39.509 20.271 1.00 12.69 AG ATOM 3030 C4 WAY 169 79.504 39.509 20.505 1.00 31.88 AG ATOM 3030 | MOTA | 1504 | OT GLY | 164 | 93.353 | 30.250 | 9.858 | 1.00 21.99 | A_13 |
| ATOM 3012 CA CA 165 73.319 39.377 1.854 1.00 40.73 AIC ATOM 3017 C5 WAY 169 67.109 35.99 20.267 1.00 38.86 A65 ATOM 3018 CF1 WAY 169 66.626 35.606 19.161 1.00 30.96 A65 ATOM 3020 C2 WAY 169 68.561 35.600 17.901 1.00 41.17 A65 ATOM 3021 C3 WAY 169 68.361 35.603 17.728 1.00 35.73 A65 ATOM 3021 C3 WAY 169 68.361 35.623 17.728 1.00 33.71 A65 ATOM 3022 C4 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3022 C4 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3023 N20 WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3024 CD WAY 169 68.861 35.862 1.00 21.141 1.00 33.16 A65 ATOM 3025 C23 WAY 169 68.861 35.862 21.141 1.00 33.16 A65 ATOM 3026 C28 WAY 169 68.861 35.862 22.685 1.00 25.69 A65 ATOM 3026 C28 WAY 169 68.986 34.739 22.685 1.00 33.61 A65 ATOM 3027 C27 WAY 169 66.91 33.061 23.490 1.00 32.16 A65 ATOM 3028 CM WAY 169 66.91 33.061 23.490 1.00 32.16 A65 ATOM 3028 CM WAY 169 66.91 33.061 23.490 1.00 32.16 A65 ATOM 3028 CM WAY 169 67.141 34.238 24.205 1.00 33.61 A65 ATOM 3029 N25 WAY 169 67.703 32.748 22.426 1.00 42.39 A65 ATOM 3030 C24 WAY 169 67.703 32.748 22.426 1.00 42.39 A65 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3031 S21 WAY 169 77.513 38.570 21.438 1.00 29.69 A66 ATOM 3033 C21 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3035 C18 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3030 C24 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3030 C36 WAY 169 73.600 39.453 20.169 1.00 11.82 A66 ATOM 3030 C38 WAY 169 73.600 39.453 20.169 1.00 11.82 A66 ATOM 3030 C38 WAY 169 73.600 39.453 20.169 1.00 11.82 A66 ATOM 3030 C38 WAY 169 73.600 39.453 20.169 1.00 11.82 A66 ATOM 3030 C38 WAY 169 73.600 39.453 20.169 1.00 11.82 A66 ATOM 3030 C38 WAY 169 73.600 39.453 20.169 1.00 13.88 A66 ATOM 3030 C38 WAY 169 73.600 39.932 20.557 1.00 26.57 A66 ATOM 3030 C38 WAY 169 73.600 39.932 20.557 1.00 26.57 A66 ATOM 3030 C38 WAY 169 73.600 39.932 20.557 1.00 26.57 A66 ATOM 3030 C38 WAY 169 73.600 39.932 20.557 1.00 22.56 BACT WAY 169 73.600 39.932 20.55 | | | | | | | | | AION AION |
| ATOM 3018 CF1 WAY 169 67.400 35.999 20.267 1.00 38.86 A65 ATOM 3019 CH WAY 169 66.26 35.606 19.161 1.00 30.96 A65 ATOM 3020 CZ WAY 169 68.501 35.623 17.728 1.00 36.26 A65 ATOM 3021 C3 WAY 169 69.339 36.039 18.811 1.00 35.73 A65 ATOM 3022 C4 WAY 169 69.339 36.039 18.811 1.00 33.71 A65 ATOM 3023 N20 WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3024 CD WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3025 C23 WAY 169 68.807 35.623 7.728 1.00 29.78 A66 ATOM 3025 C23 WAY 169 68.807 35.608 23.798 1.00 29.78 A66 ATOM 3025 C23 WAY 169 68.807 35.088 23.798 1.00 31.72 A66 ATOM 3026 C28 WAY 169 68.807 35.088 23.798 1.00 31.72 A66 ATOM 3027 C27 WAY 169 67.141 34.238 24.205 1.00 33.61 A66 ATOM 3028 CM WAY 169 67.141 34.238 24.205 1.00 33.61 A66 ATOM 3028 CW WAY 169 67.703 32.748 22.426 1.00 42.39 A66 ATOM 3030 C24 WAY 169 67.703 32.748 22.426 1.00 24.23 A66 ATOM 3030 C24 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3032 C16 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3033 C21 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3033 C21 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3030 C24 WAY 169 73.400 39.453 20.169 1.00 19.32 A66 ATOM 3031 C17 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3030 C24 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3031 C17 WAY 169 73.282 38.272 22.507 1.00 26.57 A66 ATOM 3030 C34 WAY 169 73.480 38.564 22.402 1.00 11.88 A66 ATOM 3031 C18 WAY 169 73.483 38.564 22.402 1.00 11.88 A66 ATOM 3030 C34 WAY 169 73.483 38.564 22.402 1.00 11.88 A66 ATOM 3030 C34 WAY 169 73.483 38.564 22.402 1.00 11.88 A66 ATOM 3030 C36 WAY 169 73.748 38.564 22.402 1.00 11.82 A66 ATOM 3030 C36 WAY 169 73.748 38.564 22.402 1.00 11.89 A66 ATOM 3030 C36 WAY 169 75.623 39.455 21.411 1.00 16.99 A66 ATOM 3040 O15 WAY 169 76.623 79.95 50.92 22.71 1.00 12.69 A66 ATOM 3040 O15 WAY 169 76.623 79.95 50.92 22.71 1.00 12.69 A66 ATOM 3040 O15 WAY 169 76.623 79.95 50.92 22.71 1.00 12.69 A66 ATOM 3040 O15 WAY 169 76.623 79.95 50.92 20.657 1.00 26.57 A68 ATOM 304 | | | | | | | | 1.00 11.76 | MOIA |
| ATOM 3019 CH WAY 169 67.199 35.400 17.901 1.00 41.17 A65 ATOM 3020 C2 WAY 169 68.851 35.623 17.728 1.00 35.73 A65 ATOM 3021 C3 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3022 C4 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3023 N20 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3023 N20 WAY 169 68.807 36.617 21.141 1.00 33.16 A66 ATOM 3025 C23 WAY 169 68.86 34.739 22.685 1.00 25.69 A66 ATOM 3026 C28 WAY 169 68.867 36.617 21.141 1.00 33.16 A66 ATOM 3026 C28 WAY 169 68.187 35.088 23.798 1.00 31.72 A66 ATOM 3027 C27 WAY 169 66.921 33.061 23.490 1.00 32.16 A66 ATOM 3028 CM WAY 169 66.921 33.061 23.490 1.00 32.16 A66 ATOM 3029 N25 WAY 169 66.921 33.061 23.490 1.00 32.16 A66 ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 27.88 A66 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A66 ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A66 ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A66 ATOM 3035 C19 WAY 169 72.323 39.163 20.269 1.00 19.32 A66 ATOM 3036 C18 WAY 169 72.323 39.453 20.169 1.00 11.82 A66 ATOM 3037 C17 WAY 169 72.323 39.453 20.169 1.00 11.82 A66 ATOM 3038 C30 WAY 169 75.623 39.456 21.241 1.00 16.99 A66 ATOM 3030 C24 WAY 169 77.382 38.272 22.271 1.00 26.57 A6 ATOM 3030 C34 WAY 169 77.382 38.272 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 77.382 38.272 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 77.382 38.272 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 77.582 39.445 21.141 1.00 16.99 A66 ATOM 3040 C15 WAY 169 77.582 39.445 21.141 1.00 16.99 A66 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 169 77.780 36.256 18.621 1.00 30.48 A6 ATOM 3040 C15 WAY 36 30.380 30 | MOTA | 3017 | C5 WAY | 169 | 67.400 | 35.999 | 20.267 | 1.00 38.86 | A693 |
| ATOM 3021 C3 WAY 169 68.561 35.623 17.728 1.00 36.26 A55 ATOM 3022 C4 WAY 169 69.339 36.039 18.811 1.00 35.73 A65 ATOM 3023 N20 WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3023 N20 WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3024 CD WAY 169 69.699 36.617 21.141 1.00 33.16 A65 ATOM 3025 C23 WAY 169 68.986 34.739 22.685 1.00 29.78 A65 ATOM 3026 C28 WAY 169 68.986 34.739 22.685 1.00 25.69 A65 ATOM 3026 C28 WAY 169 67.141 34.238 24.205 1.00 31.72 A65 ATOM 3028 CM WAY 169 67.703 32.748 22.426 1.00 31.61 A65 ATOM 3029 N25 WAY 169 66.921 33.061 23.490 1.00 32.16 A65 ATOM 3029 N25 WAY 169 67.703 32.748 22.426 1.00 42.39 A66 ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 27.88 A66 ATOM 3031 S21 WAY 169 67.703 32.748 22.426 1.00 42.39 A66 ATOM 3032 C16 WAY 169 73.33 38.570 21.438 1.00 29.69 A66 ATOM 3033 C21 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3035 C19 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3037 C17 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3037 C17 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.483 85.644 22.2402 1.00 11.88 A66 ATOM 3037 C17 WAY 169 72.322 38.272 22.507 1.00 26.57 A66 ATOM 3040 C15 WAY 169 75.623 39.445 21.141 1.00 19.50 A66 ATOM 3040 C15 WAY 169 70.780 39.452 21.141 1.00 19.50 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3055 CB THR 7 40.433 57.507 1.00 39.45 A66 ATOM 3040 C15 WAY 169 70.780 39.509 22.271 1.00 12.69 A66 ATOM 3054 CB WAY 169 70.780 39 | | | | | | | | | A693 A693 |
| ATOM 3022 C4 WAY 169 68.807 36.216 20.078 1.00 33.71 A65 ATOM 3023 N20 WAY 169 69.699 36.617 21.141 1.00 33.16 A69 ATOM 3024 CD WAY 169 70.137 35.640 22.189 1.00 29.78 A69 ATOM 3026 C28 WAY 169 68.986 34.739 22.685 1.00 25.69 A65 ATOM 3027 C27 WAY 169 68.986 34.739 22.685 1.00 31.72 A66 ATOM 3028 CM WAY 169 67.141 34.238 24.205 1.00 31.72 A66 ATOM 3028 CM WAY 169 67.141 34.238 24.205 1.00 32.16 A66 ATOM 3029 N25 WAY 169 67.703 32.748 22.426 1.00 42.39 A60 ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 22.18 A66 ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 24.43 A66 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A66 ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A66 ATOM 3033 C21 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.748 38.564 22.401 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.748 38.564 22.401 1.00 11.88 A66 ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A66 ATOM 3038 033 WAY 169 73.748 38.564 22.021 1.00 11.88 A66 ATOM 3030 C36 WAY 169 73.748 38.564 22.021 1.00 11.88 A66 ATOM 3030 C36 WAY 169 73.748 38.564 22.021 1.00 11.88 A66 ATOM 3030 C36 WAY 169 73.748 38.564 22.021 1.00 11.89 A66 ATOM 3030 C36 WAY 169 72.382 38.272 22.507 1.00 26.57 A66 ATOM 3034 0.015 WAY 169 72.382 38.272 22.507 1.00 26.57 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 0.15 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3050 CC THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 5050 CC THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1510 C THR 7 41.453 59.582 7.908 1.00 36.97 B_ ATOM 1520 CC LEU 8 39.406 | ATOM | 3020 | C2 WAY | 169 | 68.561 | 35.623 | 17.728 | 1.00 36.26 | A693 |
| ATOM 3024 CD WAY 169 | | | | | | | | | A693 A693 |
| ATOM 3026 C28 WAY 169 68.986 34.739 22.685 1.00 25.69 A6. ATOM 3026 C28 WAY 169 68.187 35.088 23.798 1.00 31.72 A6. ATOM 3027 C27 WAY 169 67.141 34.238 24.205 1.00 33.61 A6. ATOM 3028 NC WAY 169 66.921 33.061 23.490 1.00 32.16 A6. ATOM 3029 NC5 WAY 169 66.921 33.061 23.490 1.00 32.16 A6. ATOM 3030 C24 WAY 169 68.709 33.546 22.426 1.00 42.39 A6. ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A6. ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A6. ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A6. ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A6. ATOM 3035 C19 WAY 169 74.267 39.156 21.241 1.00 19.50 A6. ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.82 A6. ATOM 3037 C17 WAY 169 72.323 39.453 21.241 1.00 19.50 A6. ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 19.50 A6. ATOM 3039 C36 WAY 169 75.623 39.445 21.141 1.00 12.69 A6. ATOM 3030 030 WAY 169 75.623 39.445 21.141 1.00 12.69 A6. ATOM 3030 030 WAY 169 75.623 39.445 21.141 1.00 12.69 A6. ATOM 3040 015 WAY 169 69.30 39.032 20.657 1.00 13.98 A6. ATOM 3041 044 WAY 169 69.419 38.338 22.22.27 1.00 12.69 A6. ATOM 3043 NP WAY 169 77.780 36.256 18.621 1.00 12.69 A6. ATOM 3040 010 WAY 169 77.780 36.256 18.621 1.00 30.48 A6. ATOM 3040 010 WAY 169 77.780 36.256 18.621 1.00 12.69 A6. ATOM 3040 010 WAY 169 77.780 36.926 17.553 1.00 10.00 A6. ATOM 3040 010 WAY 169 77.561 36.926 17.553 1.00 10.00 A6. ATOM 3040 010 WAY 169 77.604 36.256 18.621 1.00 22.94 A6. ATOM 3040 02 WAY 169 77.604 36.256 18.621 1.00 22.94 A6. ATOM 3040 02 WAY 169 77.604 36.256 18.621 1.00 22.94 A6. ATOM 3040 03 WAY 169 77.606 60.584 36.17.575 1.00 21.20 B. ATOM 1508 C62 THR 7 40.920 59.113 6.901 1.00 30.48 A6. ATOM 3040 08 WAY 169 77.606 60.584 36.17.27 77.426 1.00 30.48 A6. ATOM 3040 08 WAY 169 77.606 60.584 36.17.27 17.426 1.00 22.94 A6. ATOM 1508 C62 THR 7 40.920 59.113 6.901 1.00 32.45 B. ATOM 1508 C62 THR 7 40.920 59.113 6.901 1.00 32.45 B. ATOM 1508 C62 THR 7 40.920 60.864 7.503 1.00 21.20 B. ATOM 1510 O THR 7 41.386 56.986 60.887 7.908 1.00 30.91 B. ATOM 1520 C6 LEU | MOTA | 3023 | | 7 169 | 69.699 | 36.617 | 21.141 | 1.00 33.16 | A693 |
| ATOM 3027 C27 WAY 169 67.141 34.238 24.205 1.00 33.61 A61 ATOM 3028 CM WAY 169 66.921 33.061 23.490 1.00 32.16 ATOM 3029 N25 WAY 169 67.703 32.748 22.426 1.00 42.39 A66 ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 27.88 A61 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A61 ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A61 ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A61 ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3035 C19 WAY 169 73.400 39.453 20.169 1.00 11.82 A66 ATOM 3036 C18 WAY 169 73.748 38.554 22.402 1.00 11.88 A61 ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A66 ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 19.50 A66 ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A66 ATOM 3040 O15 WAY 169 69.030 39.032 20.657 1.00 13.98 A66 ATOM 3041 O14 WAY 169 69.419 38.338 22.942 1.00 22.94 A61 ATOM 3044 O10 WAY 169 70.780 36.256 18.621 1.00 30.48 A62 ATOM 3044 O10 WAY 169 70.780 36.256 18.621 1.00 30.48 A62 ATOM 3044 O10 WAY 169 71.192 36.946 17.553 1.00 10.00 A62 ATOM 3045 O8 WAY 169 71.192 36.946 17.553 1.00 10.00 A62 ATOM 3045 O8 WAY 169 71.614 35.847 19.414 1.00 39.46 A63 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1506 OG1 THR 7 41.453 59.582 7.908 1.00 32.45 B_ ATOM 1507 C THR 7 40.920 59.113 6.901 1.00 33.412 B_ ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 32.45 B_ ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 32.45 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.15 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 21.78 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 21.78 B_ ATOM 1520 CG LEU 8 36.6860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.696 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.6860 61.484 6.863 1.00 27.13 B_ ATOM 1525 C THR 7 41.677 62.667 6.597 1.00 19.05 B_ ATOM 1520 CG LEU 8 36.6860 61.484 6.863 1.00 27.16 B_ ATOM 1521 CD1 LEU 8 36.696 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.696 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.662 61.866 7.298 1.00 27.16 B_ ATOM 1522 CD2 LEU 8 36.662 61.864 41.43 91 | | | | | | | | | A693 A693 |
| ATOM 3028 CM. WAY 169 66.921 33.061 23.490 1.00 32.16 A69 ATOM 3029 N25 WAY 169 67.703 32.748 22.426 1.00 42.39 AFOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 27.88 A69 ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 AFOM 3032 C16 WAY 169 72.032 39.163 20.269 1.00 19.32 A69 ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A69 ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 19.32 A69 ATOM 3035 C19 WAY 169 74.267 39.156 21.241 1.00 19.50 A69 ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A69 ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A69 ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A69 ATOM 3030 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A69 ATOM 3040 O15 WAY 169 69.419 38.338 22.942 1.00 22.94 A69 ATOM 3041 O14 WAY 169 69.419 38.338 22.942 1.00 30.48 A69 ATOM 3040 O15 WAY 169 70.780 36.256 18.621 1.00 30.48 A69 ATOM 3040 O15 WAY 169 70.780 36.256 18.621 1.00 30.48 A69 ATOM 3040 O16 WAY 169 70.780 36.256 18.621 1.00 30.48 A69 ATOM 3040 O16 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O16 WAY 169 72.581 37.127 17.426 1.00 38.25 A69 ATOM 3040 O16 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O16 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O17 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O18 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O18 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O18 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 3040 O18 WAY 169 72.581 37.127 17.426 1.00 39.46 A69 ATOM 1506 OGI THR 7 40.443 57.305 5.225 1.00 22.94 B6 ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B2 ATOM 1508 CG2 THR 7 41.4386 56.786 7.488 1.00 34.12 B2 ATOM 1510 O THR 7 41.4386 56.786 7.488 1.00 34.12 B2 ATOM 1510 O THR 7 41.4386 56.786 7.488 1.00 34.12 B2 ATOM 1510 O THR 7 41.386 56.786 7.488 1.00 27.13 B2 ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B2 ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B2 ATOM 1520 CG LEU 8 36.662 66.854 5.510 0.00 19.05 B2 ATOM 1522 CD LEU 8 40.432 61.896 7.298 1.00 27.16 B2 ATOM 1522 CD | | | | | | | | | A693 A693 |
| ATOM 3030 C24 WAY 169 68.709 33.546 22.016 1.00 27.88 A6. ATOM 3031 S21 WAY 169 69.757 38.233 21.577 1.00 24.43 A6. ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A6. ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A6. ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 11.82 A6. ATOM 3035 C19 WAY 169 74.267 39.156 21.241 1.00 19.50 A6. ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A6. ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A6. ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6. ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6. ATOM 3040 015 WAY 169 69.419 38.338 22.942 1.00 22.94 A6. ATOM 3040 015 WAY 169 69.419 38.338 22.942 1.00 22.94 A6. ATOM 3043 N9 WAY 169 70.780 36.256 18.621 1.00 30.48 A6. ATOM 3044 010 WAY 169 71.192 36.946 17.553 1.00 10.00 A6. ATOM 3046 C29 WAY 169 71.192 36.946 17.553 1.00 10.00 A6. ATOM 3046 C29 WAY 169 72.581 37.127 17.426 1.00 38.25 A6. ATOM 3046 C29 WAY 169 76.504 39.509 5.762 1.00 22.20 B. ATOM 3046 C29 WAY 169 76.504 39.509 5.762 1.00 23.15 B. ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B. ATOM 1506 CG1 THR 7 39.149 56.999 5.762 1.00 23.15 B. ATOM 1508 CC2 THR 7 40.920 59.113 6.901 1.00 32.45 B. ATOM 1508 CC2 THR 7 41.371 57.761 6.365 1.00 23.15 B. ATOM 1508 CC2 THR 7 41.371 57.761 6.365 1.00 23.60 B. ATOM 1513 N THR 7 41.386 56.786 7.908 1.00 34.12 B. ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 22.66 B. ATOM 1518 CA LEU 8 39.907 59.694 6.265 1.00 23.60 B. ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B. ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B. ATOM 1521 CD LEU 8 36.860 61.848 7.503 1.00 21.78 B. ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B. ATOM 1523 C LEU 8 36.860 61.484 6.863 1.00 27.13 B. ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B. ATOM 1523 C LEU 8 36.626 61.894 6.669 1.00 46.24 B. ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.16 B. ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B. ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B. ATOM 1524 C D LEU 8 64.4143 9.148 1.00 32.24 B. ATO | MOTA | 3028 | CM. WAY | 169 | 66.921 | 33.061 | 23.490 | 1.00 32.16 | A693 |
| ATOM 3031 S21 WAY 169 69.757 38.213 21.577 1.00 24.43 A6. ATOM 3032 C16 WAY 169 71.513 38.570 21.438 1.00 29.69 A6. ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A6. ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 11.82 A6. ATOM 3035 C19 WAY 169 73.400 39.453 20.169 1.00 11.82 A6. ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A6. ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A6. ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6. ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6. ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6. ATOM 3040 015 WAY 169 69.030 39.032 20.657 1.00 13.98 A6. ATOM 3041 014 WAY 169 69.419 38.338 22.942 1.00 22.94 A6. ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6. ATOM 3044 010 WAY 169 70.780 36.256 18.621 1.00 30.48 A6. ATOM 3044 010 WAY 169 71.192 36.946 17.553 1.00 10.00 A6. ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6. ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 39.46 A6. ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 39.46 A6. ATOM 3046 C29 WAY 169 72.581 37.127 17.426 1.00 39.46 A6. ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6. ATOM 1505 CB THR 7 40.443 57.305 5.255 1.00 21.20 B_ ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ ATOM 1510 O THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1511 CH R 7 41.371 57.761 6.365 1.00 22.66 B_ ATOM 1512 CD1 LEU 8 39.397 60.984 6.669 1.00 22.66 B_ ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 22.66 B_ ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.23 B_ ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 21.78 B_ ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1522 CD LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.0 | | | | | | | | | A693 A693 |
| ATOM 3033 C21 WAY 169 72.032 39.163 20.269 1.00 19.32 A6. ATOM 3034 C20 WAY 169 73.400 39.453 20.169 1.00 11.82 A6. ATOM 3035 C19 WAY 169 74.267 39.156 21.241 1.00 19.50 A6. ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A6. ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A6. ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6. ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6. ATOM 3040 015 WAY 169 69.030 39.032 20.657 1.00 13.98 A6. ATOM 3041 014 WAY 169 69.419 38.338 22.942 1.00 22.94 A6. ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6. ATOM 3044 010 WAY 169 70.780 36.256 18.621 1.00 30.48 A6. ATOM 3044 010 WAY 169 71.192 36.946 17.553 1.00 10.00 A6. ATOM 3045 08 WAY 169 71.192 36.946 17.553 1.00 10.00 A6. ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6. ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6. ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 21.20 B_ATOM 1506 OG1 THR 7 40.443 57.305 5.225 1.00 21.20 B_ATOM 1506 OG1 THR 7 40.435 59.582 7.908 1.00 32.45 B_ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ATOM 1510 O THR 7 41.386 56.786 7.488 1.00 34.12 B_ATOM 1510 O THR 7 41.386 56.786 7.488 1.00 34.12 B_ATOM 1510 CB LEU 8 39.907 59.694 6.265 1.00 22.66 B_ATOM 1510 CB LEU 8 39.907 59.694 6.265 1.00 22.66 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.18 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.18 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.18 B_ATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.18 B_ATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.18 B_ATOM 1522 CD2 LEU 8 36.860 61.484 6.861 1.00 27.84 B_ATOM 1525 CD LEU 8 36.622 60.854 5.510 1.00 19.23 B_ATOM 1525 CD LEU 8 36.622 60.854 5.510 1.00 19.23 B_ATOM 1522 CD2 LEU 8 36.662 60.854 5.510 1.00 19.23 B_ATOM 1522 CD2 LEU 8 36.662 61.804 8.618 1.00 27.84 B_ATOM 1525 CD LEU 8 36.662 60.854 5.510 1.00 19.23 B_ATOM 1522 CD2 LEU 8 41.077 62.667 6.597 1.00 46.24 B_ATOM 1525 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ATOM 1528 CB LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1528 CB LYS 9 40.615 | MOTA | 3031 | S21 WAY | Y 169 | 69.757 | 38.213 | 21.577 | 1.00 24.43 | A693 |
| ATOM 3035 C19 WAY 169 74.267 39.156 21.241 1.00 19.50 A6 ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A6 ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A6 ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6 ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6 ATOM 3040 015 WAY 169 69.030 39.032 20.657 1.00 13.98 A6 ATOM 3041 014 WAY 169 69.419 38.338 22.942 1.00 22.94 A6 ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6 ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 39.46 A6 ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ ATOM 1508 CG2 THR 7 40.920 59.113 6.901 1.00 32.45 B_ ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ ATOM 1510 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1515 CA THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 22.66 B_ ATOM 1518 CA LEU 8 39.907 59.694 6.265 1.00 22.66 B_ ATOM 1521 CD1 LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1522 CD2 LEU 8 36.862 60.854 5.510 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.860 61.894 7.503 1.00 19.05 B_ ATOM 1524 CD1 LEU 8 36.860 61.896 7.298 1.00 27.16 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.18 B_ ATOM 1522 CD2 LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1524 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.18 B_ ATOM 1522 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1524 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.16 B_ ATOM 1524 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1524 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 CD LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ | | | | | | | | | A693 A693 |
| ATOM 3036 C18 WAY 169 73.748 38.564 22.402 1.00 11.88 A6 ATOM 3037 C17 WAY 169 72.382 38.272 22.507 1.00 26.57 A6 ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6 ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6 ATOM 3040 O15 WAY 169 69.030 39.032 20.657 1.00 13.98 A6 ATOM 3041 O14 WAY 169 69.419 38.338 22.942 1.00 22.94 A6 ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6 ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 O10 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3044 O10 WAY 169 72.581 37.127 17.426 1.00 39.46 A6 ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1506 OG1 THR 7 40.943 57.305 5.225 1.00 21.20 B_ ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ ATOM 1516 N LEU 8 39.387 60.984 6.649 1.00 22.66 B_ ATOM 1516 N LEU 8 39.387 60.984 6.649 1.00 22.66 B_ ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B_ ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1524 O LEU 8 40.432 61.896 7.298 1.00 27.13 B_ ATOM 1524 CD2 LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 CD2 LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1526 CB LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1526 CB LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1526 CB LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1527 CA LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | | | | | | | | | A693 |
| ATOM 3038 033 WAY 169 75.623 39.445 21.141 1.00 16.99 A6 ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6 ATOM 3040 015 WAY 169 69.030 39.032 20.657 1.00 13.98 A6 ATOM 3041 014 WAY 169 69.419 38.338 22.942 1.00 22.94 A6 ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6 ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3045 08 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 BATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 BATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 BATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 BATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 BATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 BATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 BATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 BATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 BATOM 1520 CG LEU 8 38.113 60.848 7.503 1.00 21.78 BATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.13 BATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.13 BATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.05 BATOM 1524 O LEU 8 40.432 61.896 67.298 1.00 27.16 BATOM 1524 O LEU 8 40.432 61.896 67.298 1.00 27.16 BATOM 1527 CA LYS 9 40.615 61.804 8.618 1.00 27.16 BATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 17.84 BATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 BATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | MOTA | 3036 | C18 WA | Y 169 | 73.748 | 38.564 | 22.402 | 1.00 11.88 | A693 |
| ATOM 3039 C36 WAY 169 76.504 39.509 22.271 1.00 12.69 A6 ATOM 3040 015 WAY 169 69.030 39.032 20.657 1.00 13.98 A6 ATOM 3041 014 WAY 169 69.419 38.338 29.42 1.00 22.94 A6 ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6 ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3046 C29 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 32.45 B_ ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1516 N LEU 8 39.907 59.694 6.649 1.00 22.666 B_ ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.666 B_ ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B_ ATOM 1524 O LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1526 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | | | | | | | | | A693 A693 |
| ATOM 3041 014 WAY 169 69.419 38.338 22.942 1.00 22.94 A6 ATOM 3042 C7 WAY 169 70.780 36.256 18.621 1.00 30.48 A6 ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3045 08 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B_ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ATOM 1522 CD2 LEU 8 36.860 61.484 6.863 1.00 27.13 B_ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ATOM 1524 O LEU 8 40.432 61.896 7.298 1.00 27.16 B_ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1525 CA LYS 9 41.572 62.667 6.597 1.00 46.24 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 27.84 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 27.84 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 27.84 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 27.84 B_ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 27.84 B_ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | MOTA | 3039 | C36 WA | Y 169 | 76.504 | 39.509 | 22.271 | 1.00 12.69 | A693 |
| ATOM 3043 N9 WAY 169 71.192 36.946 17.553 1.00 10.00 A6 ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3045 08 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B ATOM 1527 CA LYS 9 40.615 61.804 8.618 1.00 27.18 B ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B | | | · · · - | | | | | | A693 A693 |
| ATOM 3044 010 WAY 169 72.581 37.127 17.426 1.00 38.25 A6 ATOM 3045 08 WAY 169 71.614 35.847 19.414 1.00 39.46 A6 ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ ATOM 1510 O THR 7 40.920 59.113 6.901 1.00 32.45 B_ ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B_ ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B_ ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1526 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | | | | | | | | | A693 A693 |
| ATOM 3046 C29 WAY 169 66.584 36.175 21.566 1.00 46.13 A6 ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 39.663 64.342 8.853 1.00 29.47 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | MOTA | 3044 | 010 WA | Y 169 | 72.581 | . 37.127 | 17.426 | 1.00 38.25 | A693 |
| ATOM 1505 CB THR 7 40.443 57.305 5.225 1.00 21.20 B_ATOM 1506 OG1 THR 7 39.149 56.999 5.762 1.00 25.31 B_ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B_ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B_ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ATOM 1522 CD2 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1527 CA LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1527 CA LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | | | | | | | | | A693 A693 |
| ATOM 1508 CG2 THR 7 41.017 56.087 4.541 1.00 23.15 B_ATOM 1509 C THR 7 40.920 59.113 6.901 1.00 32.45 B_ATOM 1510 O THR 7 41.453 59.582 7.908 1.00 36.97 B_ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B_ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B_ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B_ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B_ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B_ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B_ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ATOM 1528 CB LYS 9 39.663 64.342 8.853 1.00 29.47 B_ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 | | | | R 7 | | 57.305 | 5.225 | 1.00 21.20 | B_13 |
| ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 39.663 64.342 8.853 1.00 29.47 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B | MOTA | 1508 | | R 7 | 41.017 | 56.087 | | 1.00 23.15 | B_13 B_13 |
| ATOM 1513 N THR 7 41.386 56.786 7.488 1.00 34.12 B ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 39.663 64.342 8.853 1.00 29.47 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B | | | | R 7 | | | | | B_13 B_13 |
| ATOM 1515 CA THR 7 41.371 57.761 6.365 1.00 26.16 B_ ATOM 1516 N LEU 8 39.907 59.694 6.265 1.00 23.60 B_ ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 B_ ATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 B_ ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B_ ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B_ ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B_ | MOTA | 1513 | N TH | R 7 | 41.386 | 56.786 | 7.488 | 1.00 34.12 | B_13 |
| ATOM 1518 CA LEU 8 39.387 60.984 6.649 1.00 22.66 BATOM 1519 CB LEU 8 38.113 60.848 7.503 1.00 21.78 BATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 BATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 BATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 BATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 BATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 BATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 BATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 BATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 BATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 BATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 BATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 BATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 | | | | R 7 | | | | | B_13 B_13 |
| ATOM 1520 CG LEU 8 36.860 61.484 6.863 1.00 27.13 B_ ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 B_ ATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 B_ ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B_ ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B_ ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B_ | MOTA | 1518 | CA LE | U 8 | 39.387 | 60.984 | 6.649 | 1.00 22.66 | B_13 |
| ATOM 1521 CD1 LEU 8 36.996 63.016 6.705 1.00 19.05 BATOM 1522 CD2 LEU 8 36.622 60.854 5.510 1.00 19.23 BATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 BATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 BATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 BATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 BATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 BATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 BATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 BATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 BATOM | | | | U 8 | | | | | B_13 ·B_13 |
| ATOM 1523 C LEU 8 40.432 61.896 7.298 1.00 27.16 B ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B | | | | 8 U | | | 6.705 | 1.00 19.05 | B_13 |
| ATOM 1524 O LEU 8 41.077 62.667 6.597 1.00 46.24 B_ ATOM 1525 N LYS 9 40.615 61.804 8.618 1.00 27.84 B_ ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B_ ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B_ | MOTA | 1523 | C LE | 8 U. | 40.432 | 61.896 | | 1.00 27.16 | B_13 B_13 |
| ATOM 1527 CA LYS 9 41.572 62.674 9.306 1.00 15.20 B ATOM 1528 CB LYS 9 41.147 64.143 9.148 1.00 32.32 B ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B | | | | U 8 S 9 | | | | | B_13 B_13 |
| ATOM 1528 CB LIS 9 41.147 64.143 9.148 1.00 32.32 B_ ATOM 1529 CG LYS 9 39.663 64.342 8.853 1.00 29.47 B_ ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B_ | MOTA | 1527 | CA LY | s 9 | 41.572 | 62.674 | 9.306 | 1.00 15.20 | B_13 |
| ATOM 1530 CD LYS 9 38.788 64.243 10.084 1.00 28.34 B | MOTA | | | s 9 | | | | | B_13 B_13 |
| | MOTA | 1530 | CD LY | 'S 9 | | | | | _B_13 |

| ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM | 1531 1532 1536 1537 1538 1540 1541 1542 1543 1544 | NZ C O N CA CB CCD | LYS LYS LYS TRP TRP TRP TRP | 9 9 9 10 10 10 10 | 38.830 38.732 41.809 41.268 42.654 42.988 44.403 45.499 46.077 47.071 | 65.556 66.725 62.384 61.428 63.208 63.112 63.660 62.890 61.650 61.302 | 10.842 9.888 10.780 11.334 11.390 12.813 13.048 12.349 12.762 11.829 | 1.00 18.48 1.00 33.19 1.00 20.69 1.00 25.62 1.00 12.09 1.00 21.78 1.00 23.03 1.00 27.60 1.00 27.28 1.00 22.11 | B_13 B_13 B_13 B_13 B_13 B_13 B_13 B_13 |
|--|--|--------------------------------------|---|-------------------------------------|--|--|---|--|--|
| MOTA MOTA MOTA | 1545 1546 1547 | CD1 | TRP TRP | 10 10 10 | 45.859 46.153 47.094 | 60.781 63.247 62.305 | 13.847 11.198 | 1.00 11.66 1.00 21.84 | B_13 B_13 |
| MOTA MOTA | 1549 1550 | CZ2 CZ3 | TRP TRP | 10 10 | 47.847 46.632 | 60.143 59.622 | 10.873 11.929 13.951 | 1.00 10.00 1.00 25.24 1.00 22.71 | B_13 B_13 B_13 |
| MOTA MOTA | 1551 1552 | CH2 | TRP | 10 10 | 47.611 41.987 | 59.317 63.915 | 12.999 13.679 | 1.00 15.23 1.00 30.88 | B_13 B_13 |
| MOTA | 1553 | 0 | TRP | 10 | 41.673 | 65.062 | 13.359 | 1.00 32.03 | B_13 |
| MOTA MOTA | 1554 1556 | N CA | SER SER | 11 11 | 41.495 40.548 | 63.316 63.981 | 14.765 15.665 | 1.00 35.64 1.00 30.37 | B_13 |
| MOTA | 1557 | CB | SER | 11 | 39.498 | 62.995 | 16.176 | 1.00 30.37 | B_13 B_13 |
| MOTA MOTA | 1558 1560 | OG C | SER SER | 11 11 | 38.485 41.206 | 62.815 64.691 | 15.202 16.840 | 1.00 41.11 1.00 20.70 | B_13 |
| MOTA | 1561 | 0 | SER | 11 | 40.558 | 65.002 | 17.838 | 1.00 26.70 | B_13 B_13 |
| ATOM ATOM | 1562 1564 | N CA | LYS LYS | 12 12 | 42.504 43.257 | 64.910 65.607 | 16.731 | 1.00 23.56 | B_13 |
| MOTA | 1565 | CB | LYS | 12 | 43.257 | 64.631 | 17.756 18.688 | 1.00 15.00 1.00 18.58 | B_13 B_13 |
| MOTA MOTA | 1566 1567 | CG CD | LYS LYS | 12 12 | 44.658 | 63.452 | 18.010 | 1.00 15.94 | B_13 |
| MOTA | 1568 | CE | LYS | 12 | 45.456 | 62.589 61.715 | .19.007 19.933 | 1.00 23.03 1.00 27.10 | B_13 B_13 |
| MOTA MOTA | 1569 | NZ | LYS | 12 | 44.075 | 62.402 | 21.157 | 1.00 34.75 | B_13 |
| MOTA | 1573 1574 | С О | LYS | 12 12 | 44.200 44.567 | 66.453 66.039 | 16.914 15.808 | 1.00 25.03 1.00 25.20 | B_13 B_13 |
| MOTA | 1575 | N | MET | 13 | 44.536 | 67.647 | 17.401 | 1.00 18.44 | B_13 |
| MOTA MOTA | 1577 1578 | CA CB | MET MET | 13 13 | 45.377 44.864 | 68.582 70.015 | 16.663 16.880 | 1.00 24.63 1.00 13.15 | B_13 B_13 |
| MOTA | 1579 | CG | MET | 13 | 43.421 | 70.253 | 16.419 | 1.00 21.56 | B_13 |
| ATOM ATOM | 1580 1581 | SD CE | MET MET | 13 13 | 43.167 41.433 | 70.131 69.678 | 14.616 14.474 | 1.00 31.39 1.00 24.70 | B_13 B_13 |
| MOTA | 1582 | C | MET | 13 | 46.850 | 68.468 | 17.034 | 1.00 11.65 | B_13 |
| ATOM ATOM | 1583 1584 | O N | MET ASN | 13 14 | 47.728 47.103 | 68.815 67.985 | 16.247 18.242 | 1.00 14.33 1.00 16.99 | B_13 B_13 |
| MOTA | 1586 | CA | ASN | 14 | 48.448 | 67.793 | 18.760 | 1.00 24.42 | B_13 B_13 |
| MOTA MOTA | 1587 1588 | CB CG | ASN ASN | 14 14 | 48.437 47.896 | 68.006 69.356 | 20.268 20.633 | 1.00 17.84 1.00 35.10 | B_13 B_13 |
| MOTA | 1589 | OD1 | ASN | 14 | 48.614 | 70.346 | 20.560 | 1.00 34.88 | B_13 |
| MOTA MOTA | 1590 1593 | ND2 C | ASN ASN | 14 14 | 46.610 48.831 | 69.424 66.364 | 20.955 18.421 | 1.00 32.98 1.00 22.70 | B_13 |
| MOTA | 1594 | 0 | ASN | 14 | 48.278 | 65.405 | 18.976 | 1.00 26.03 | B_13 B_13 |
| MOTA MOTA | 1595 1597 | N· CA | LEU | 15 15 | 49.706 50.144 | 66.228 64.912 | 17.432 16.969 | 1.00 18.07 1.00 29.36 | B_13 |
| MOTA | 1598 | CB | LEU | 15 | 49.878 | 64.775 | 15.466 | 1.00 24.35 | B_13 B_13 |
| MOTA MOTA | 1599 1600 | CG CD1 | LEU | 15 15 | 48.380 48.079 | 64.762 65.469 | 15.162 13.852 | 1.00 19.51 1.00 27.59 | B_13 B_13 |
| MOTA | 1601 | CD2 | LEU | 15 | 47.902 | 63.326 | 15.163 | 1.00 19.66 | B_13 |
| MOTA MOTA | 1602 1603 | C O | LEU | 15 15 | 51.613 52.341 | 64.704 65.657 | 17.257 17.552 | 1.00 28.48 1.00 22.28 | B_13 B_13 |
| MOTA | 1604 | N | THR | 16 | 52.044 | 63.453 | 17.198 | 1.00 12.77 | B_13 |
| MOTA | 1606 1607 | CA CB | THR THR | 16 16 | 53.433 53.607 | 63.158 62.243 | 17.446 18.682 | 1.00 16.59 1.00 24.73 | B_13 |
| MOTA | 1608 | OG1 | THR | 16 | 52.912 | 61.005 | 18.481 | 1.00 24.73 | B_13 B_13 |
| MOTA MOTA | 1610 1611 | CG2 C | THR | 16 16 | 53.059 54.038 | 62.933 62.515 | 19.924 16.214 | 1.00 25.34 | B_13 |
| MOTA | 1612 | Ö | THR | | 53.315 | 62.116 | 15.214 | 1.00 21.94 1.00 19.60 | B_13 B_13 |
| MOTA MOTA | 1613 1615 | N CA | TYR TYR | | 55.365 | 62.453 | 16.184 | 1.00 18.25 | B_13 |
| MOTA | 1616 | CB | TYR | | 56.092 56.300 | 61.810 62.753 | 15.097 13.910 | 1.00 19.54 1.00 16.87 | B_13 B_13 |
| ATOM ATOM | 1617 1618 | CG CD1 | TYR | | 57.277 | 63.892 | 14.116 | 1.00 27.90 | B_13 |
| ATOM | 1619 | CD1 CE1 | | | 56.839 57.700 | 65.135 66.221 | 14.587 14.652 | 1.00 13.93 1.00 17.08 | B_13 B_13 |
| MOTA MOTA | 1620 | CD2 | TYR | 17 | 58.613 | 63.764 | 13.723 | 1.00 14.99 | B_13 |
| ATOM | 1621 1622 | CEZ | TYR TYR | | 59.479 59.017 | 64.841 66.075 | 13.777 14.242 | 1.00 25.98 1.00 33.12 | B_13 B_13 |
| MOTA MOTA | 1623 1625 | OH. | TYR | 17 | 59.866 | 67.163 | 14.276 | 1.00 23.31 | B_13 |
| MOTA | 1626 | C O | TYR TYR | | 57.417 57.895 | 61.318 61.827 | 15.650 16.668 | 1.00 18.57 1.00 26.60 | B_13 B_13 |
| ATOM | 1627 | N | ARG | 18 | 57.973 | 60.286 | 15.030 | 1.00 13.01 | B_13 |

| | | _ | | | | • | | | |
|--------|------|-----|-----|----------|--------|--------|--------|-------------|------|
| ATOM . | 1629 | CA | ARG | 18 | 59.245 | 59.750 | 15.492 | 1.00 18.74 | B_13 |
| ATOM | 1530 | CB | ARG | 18 | 59.033 | 58.589 | 16.473 | 1.00 11.96 | B_13 |
| MOTA | 1631 | CG | ARG | 18 | 60.320 | 57.911 | 16.970 | 1.00 15.06 | B_13 |
| ATOM | 1632 | CD | ARG | 18 | 60.012 | | | | |
| | | | | | | 56.596 | 17.690 | 1.00 11.72 | B_13 |
| ATOM | 1633 | NE | ARG | 18 | 61.165 | 55.689 | 17.752 | 1.00 10.00 | B_13 |
| MOTA | 1635 | CZ | ARG | 18 | 61.134 | 54.428 | 18.181 | 1.00 24.87 | B_13 |
| MOTA | 1636 | NHl | ARG | 18 | 60.004 | 53.882 | 18.614 | 1.00 13.34 | B_13 |
| ATOM | 1639 | NH2 | ARG | 18 | 62.247 | 53.703 | 18.169 | 1.00 20.03 | B_13 |
| | 1642 | C | | 18 | | | | | |
| ATOM | | | ARG | | 60.076 | 59.309 | 14.307 | 1.00 13.14 | B_13 |
| MOTA | 1643 | 0 | ARG | 18 | 59.598 | 58.588 | 13.434 | 1.00 14.10 | B_13 |
| MOTA | 1644 | N | ILE | 19 | 61.304 | 59.813 | 14.252 | 1.00 15.55 | B_13 |
| ATOM | 1646 | CA | ILE | 19 | 62.238 | 59.476 | 13.193 | 1.00 10.41 | B_13 |
| MOTA | 1647 | CB | ILE | 19 | 63.307 | 60.603 | | | |
| | | | | | | | 13.054 | 1.00 17.20 | B_13 |
| MOTA | 1648 | CG2 | | 19 | 64.273 | 60.307 | 11.903 | 1.00 16.57 | B_13 |
| MOTA | 1649 | CG1 | ILE | 19 | 62.613 | 61.952 | 12.836 | 1.00 15.47 | B_13 |
| ATOM | 1650 | CD1 | ILE | 19 | 63.543 | 63.110 | 12.783 | 1.00 14.99 | B_13 |
| MOTA | 1651 | С | ILE | 19 | 62.870 | 58.166 | 13.673 | 1.00 10.00 | B_13 |
| ATOM | 1652 | Ö | ILE | 19 | 63.829 | 58.179 | | | |
| | | - | | | | | 14.434 | 1.00 10.00 | B_13 |
| ATOM | 1653 | N | VAL | 20 | 62.289 | 57.037 | 13.276 | 1.00 17.84 | B_13 |
| ATOM | 1655 | CA | VAL | 20 | 62.785 | 55.716 | 13.696 | 1.00 16.43 | B_13 |
| ATOM | 1656 | CB | VAL | 20 | 61.911 | 54.570 | 13.138 | 1.00 13.17. | B_13 |
| ATOM | 1657 | CG1 | VAL | 20 | 62.519 | 53.208 | 13.493 | 1.00 10.00 | B_13 |
| ATOM | 1658 | CG2 | | 20 | 60.521 | 54.673 | 13.698 | 1.00 10.00 | ·— |
| | | | | | | | | | B_13 |
| ATOM | 1659 | C | VAL | 20 | 64.268 | 55.449 | 13.387 | 1.00 16.02 | B_13 |
| MOTA | 1660 | 0 | VAL | 20 | 65.001 | 54.909 | 14.218 | 1.00 21.07 | B_13 |
| ATOM | 1661 | N | asn | 21 | 64.698 | 55.762 | 12.177 | 1.00 10.00 | B_13 |
| MOTA | 1663 | CA | ASN | 21 | 66.098 | 55.571 | 11.830 | 1.00 22.13 | B_13 |
| ATOM | 1664 | CB | ASN | 21 | 66.392 | 54.128 | 11.386 | 1.00 19.75 | B_13 |
| ATOM | 1665 | CG | ASN | 21 | 65.549 | 53.673 | 10.212 | 1.00 17.63 | |
| | | | | | | | | | B_13 |
| ATOM | 1666 | | ASN | 21 | 65.329 | 52.477 | 10.042 | 1.00 31.82 | B_13 |
| MOTA | 1667 | | ASN | 21 | 65.109 | 54.602 | 9.375 | 1.00 11.42 | B_13 |
| MOTA | 1670 | С | ASN | 21 | 66.504 | 56.645 | 10.821 | 1.00 10.14 | B_13 |
| ATOM | 1671 | 0 | ASN | 21 | 65.639 | 57.377 | 10.340 | 1.00 11.74 | B_13 |
| MOTA | 1672 | N | TYR | 22 | 67.787 | 56.759 | 10.498 | 1.00 12.25 | B_13 |
| MOTA | 1674 | CA | TYR | 22 | 68.233 | 57.829 | 9.602 | 1.00 12.46 | B_13 |
| MOTA | 1675 | CB | TYR | 22 | 69.136 | 58.800 | | 1.00 23.15 | |
| | | | | | | | | | B_13 |
| MOTA | 1676 | CG | TYR | 22 | 68.461 | 59.584 | | 1.00 21.95 | B_13 |
| MOTA | 1677 | CD1 | | 22 | 68.221 | 60.945 | 11.348 | 1.00 22.29 | B_13 |
| ATOM | 1678 | CE1 | TYR | 22 | 67.625 | 61.678 | 12.347 | 1.00 10.00 | B_13 |
| MOTA | 1679 | CD2 | TYR | 22 | 68.077 | 58.974 | 12.687 | 1.00 13.42 | B_13 |
| MOTA | 1680 | CE2 | TYR | 22 | 67.471 | 59.710 | 13.693 | 1.00 14.69 | B_13 |
| MOTA | 1681 | CZ | TYR | 22 | 67.254 | 61.064 | 13.505 | 1.00 12.89 | B_13 |
| MOTA | 1682 | OH | TYR | 22 | 66.660 | 61.829 | | 1.00 16.56 | |
| | | | | | | | 14.466 | | B_13 |
| ATOM | 1684 | C | TYR | 22 | 68.988 | 57.395 | 8.359 | | B_13 |
| MOTA | 1685 | 0 | TYR | 22 | 69.793 | 56.478 | 8.407 | 1.00 16.23 | B_13 |
| MOTA | 1686 | N | THR | 23 | 68.792 | 58.111 | 7.261 | 1.00 10.39 | B_13 |
| ATOM | 1688 | CA | THR | 23 | 69.503 | 57.800 | 6.024 | 1.00 20.36 | B_13 |
| MOTA | 1689 | CB | THR | 23 | 68.909 | 58.582 | 4.829 | 1.00 16.21 | B_13 |
| MOTA | 1690 | OG1 | | 23 | 69.801 | 58.512 | 3.706 | 1.00 19.72 | |
| ATOM | 1692 | CG2 | | 23 | | | | | B_13 |
| | | | | | 68.663 | 60.039 | | | B_13 |
| MOTA | 1693 | C | THR | 23 | 70.990 | 58.153 | 6.163 | 1.00 17.35 | B_13 |
| MOTA | 1694 | 0 | THR | 23 | 71.377 | 58.958 | 7.024 | 1.00 13.88 | B_13 |
| MOTA | 1695 | N | PRO | 24 | 71.852 | 57.503 | 5.364 | 1.00 15.86 | B_13 |
| ATOM | 1696 | CD | PRO | 24 | 71.625 | 56.247 | 4.629 | 1.00 17.29 | B_13 |
| MOTA | 1697 | CA | PRO | 24 | 73.287 | 57.796 | | | B_13 |
| MOTA | 1698 | CB | PRO | 24 | 73.920 | 56.570 | | | |
| ATOM | | CG | | | | | | | B_13 |
| | 1699 | _ | PRO | 24 | 72.891 | 55.504 | | | B_13 |
| ATOM | 1700 | C | PRO | 24 | 73.635 | 59.069 | | 1.00 27.08 | B_13 |
| MOTA | 1701 | 0 | PRO | 24 | 74.698 | 59.656 | 4.869 | 1.00 19.47 | B_13 |
| MOTA | 1702 | N | ASP | 25 | 72.728 | 59.489 | 3.794 | 1.00 16.99 | B_13 |
| MOTA | 1704 | CA | ASP | 25 | 72.927 | 60.663 | | 1.00 10.00 | B_13 |
| MOTA | 1705 | CB | ASP | 25 | 71.792 | 60.758 | | | B_13 |
| ATOM | 1706 | CG | ASP | 25 | 71.665 | | | | |
| | | | | | | 59.521 | | | B_13 |
| MOTA | 1707 | OD1 | | 25 | 70.570 | 59.311 | | | B_13 |
| ATOM | 1708 | OD2 | | 25 | 72.653 | 58.762 | - | | B_13 |
| ATOM | 1709 | | ASP | 25 | 73.068 | 62.011 | 3.642 | 1.00 23.36 | B_13 |
| ATOM | 1710 | 0 | ASP | 25 | 73.694 | | | | B_13 |
| ATOM | 1711 | N | MET | 26 | 72.480 | 62.158 | | | B_13 |
| ATOM | 1713 | CA | MET | 26 | 72.510 | | | | B_13 |
| ATOM | 1714 | CB | MET | 26 | | | | | |
| | | | | | 71.154 | | | | B_13 |
| MOTA | 1715 | | MET | 26 26 | 70.782 | | | | B_13 |
| ATOM | 1716 | | MET | 26 | 69.016 | | | | B_13 |
| ATOM | 1717 | | MET | 26 | 68.395 | | | | B_13 |
| ATOM | 1718 | | MET | 26 | 72.827 | 63.238 | 7.024 | 1.00 28.80 | B_13 |
| MOTA | 1719 | 0 | MET | 26 | 72.839 | | | | B_13 |
| ATOM | 1720 | | THR | 27 | 73.157 | | | | B_13 |
| ATOM | 1722 | | THR | 27 | 73.456 | | - | | B_13 |
| | | | • | | | | | | |

| ATOM | 1723 | CB 3 | THR | 27 | 74.117 | 65.605 | 9.602 | 1.00 33.46 | B_13 |
|--------|------|-------|-----|------|--------|--------|--------|------------|--------------|
| ATOM | 1724 | OG1 1 | THR | 27 | 73.209 | 66.702 | 9.415 | 1.00 10.00 | B_13 |
| ATOM | 1726 | CG2 | THR | 27 | 75.405 | 65.863 | 8.818 | 1.00 16.30 | B_13 |
| ATOM | 1727 | | THR | 27 | 72.135 | 64.113 | 9.861 | 1.00 10.67 | |
| ATOM | 1728 | | THR | 27 | 71.072 | 64.343 | 9.281 | 1.00 16.26 | B_13 |
| ATOM | 1729 | | HIS | 28 | 72.193 | | | | B_13 |
| ATOM | 1731 | _ | | | | 63.691 | 11.124 | 1.00 18.13 | B_13 |
| | | | HIS | 28 | 70.986 | 63.514 | 11.915 | 1.00 10.00 | B_13 |
| ATOM | 1732 | | HIS | 28 | 71.322 | 63.033 | 13.333 | 1.00 10.00 | B_13 |
| ATOM | 1733 | | HIS | 28 | 71.793 | 61.608 | 13,401 | 1.00 22.65 | B_13 |
| MOTA | 1734 | CD2 1 | HIS | 28 | 72.893 | 61.003 | 12.889 | 1.00 22.73 | B_13 |
| ATOM | 1735 | ND1 | HIS | 28 | 71.103 | 60.627 | 14.080 | 1.00 19.90 | B_13 |
| ATOM | 1737 | CE1 | | 28 | 71.755 | 59.481 | 13.985 | 1.00 16.52 | |
| ATOM | 1738 | | HIS | 28 | 72.843 | 59.681 | | | B_13 |
| ATOM | 1740 | | | | | | 13.268 | 1.00 20.38 | B_13 |
| | | | HIS | 28 | 70.281 | 64.870 | 11.957 | 1.00 29.38 | B_13 |
| ATOM | 1741 | | HIS | 28 | 69.074 | 64.941 | 11.742 | 1.00 17.20 | B_13 |
| ATOM | 1742 | | SER | 29 | 71.056 | 65.944 | 12.153 | 1.00 23.96 | B_13 |
| MOTA | 1744 | CA | SER | 29 | 70.533 | 67.322 | 12.192 | 1.00 15.01 | B_13 |
| ATOM | 1745 | CB | SER | 29 | 71.661 | 68.334 | 12.438 | 1.00 14.05 | B_13 |
| ATOM | 1746 | OG | SER | 29 | 72.117 | 68.303 | 13.770 | 1.00 18.32 | B_13 |
| ATOM . | 1748 | | SER | 29 | 69.808 | 67.729 | 10.909 | 1.00 10.95 | |
| ATOM | 1749 | | SER | 29 | 68.732 | 68.314 | 10.971 | | B_13 |
| ATOM | 1750 | | GLU | 30 | 70.415 | | | 1.00 24.24 | B_13 |
| ATOM | 1752 | | | | | 67.449 | 9.757 | 1.00 10.96 | B_13 |
| | | | GLU | 30 | 69.820 | 67.786 | 8.470 | 1.00 10.00 | B_:13 |
| ATOM | 1753 | | GLU | 30 | 70.715 | 67.330 | 7.309 | 1.00 10.12 | B_13 |
| ATOM | 1754 | | GLU | 30 | 71.967 | 68.143 | 7.042 | 1.00 22.31 | B_13 |
| ATOM | 1755 | | GLU | 30 | 72.823 | 67.529 | 5.930 | 1.00 10.15 | B_13 |
| MOTA | 1756 | OE1 | GLU | 30 | 72.533 | 67.753 | 4.749 | 1.00 31.98 | B_13 |
| ATOM | 1757 | OE2 | GLU | 30 | 73.796 | 66.817 | 6.223 | 1.00 29.59 | B_13 |
| ATOM | 1758 | C | GLU | 30 | 68.481 | 67.073 | 8.336 | 1.00 20.17 | B_13 |
| MOTA | 1759 | | GLU | 30 | 67.493 | 67.685 | 7.943 | 1.00 14.31 | |
| MOTA | 1760 | | VAL | 31 | 68.451 | 65.777 | | | B_13 |
| ATOM | 1762 | | VAL | | | | 8.665 | 1.00 19.26 | B_13 |
| | | | | 31 | 67.228 | 64.989 | 8.536 | 1.00 14.22 | B_13 |
| MOTA | 1763 | | VAL | 31 | 67.472 | 63.487 | 8.716 | 1.00 17.05 | B_13 |
| MOTA | 1764 | CG1 | | 31 | 66.144 | 62.749 | 8.791 | 1.00 28.55 | B_13 |
| ATOM | 1765 | CG2 | VAL | 31 | 68.269 | 62.935 | 7.548 | 1.00 10.54 | B_13 |
| ATOM | 1766 | C | VAL | 31 | 66.138 | 65.458 | 9.477 | 1.00 12.36 | B_13 |
| ATOM | 1767 | 0 | VAL | 31 | 64.963 | 65.488 | 9.093 | 1.00 12.83 | B_13 |
| ATOM | 1768 | | GLU | 32 | 66.530 | 65.805 | 10.703 | 1.00 20.46 | |
| ATOM | 1770 | | GLU | 32 | 65.596 | 66.306 | 11.710 | 1.00 20.40 | B_13 |
| ATOM | 1771 | | GLU | 32 | | | | | B_13 |
| ATOM | 1772 | | | | 66.269 | 66.365 | 13.094 | 1.00 14.71 | B_13 |
| | | | GLU | 32 | 66.512 | 64.985 | 13.741 | 1.00 23.30 | B_13 |
| ATOM | 1773 | | GLU | 32 | 67.724 | 64.930 | 14.700 | 1.00 21.41 | B_13 |
| MOTA | 1774 | | GLU | 32 | 68.229 | 63.823 | 15.003 | 1.00 15.79 | B_13 |
| MOTA | 1775 | | GLU | 32 | 68.183 | 65.985 | 15.157 | 1.00 13.71 | B_13 |
| MOTA | 1776 | C | GLU | 32 | 65.125 | 67.697 | 11.257 | 1.00 27.19 | B_13 |
| ATOM. | 1777 | 0 | GLU | 32 | 63.951 | 68.042 | 11.383 | 1.00 19.82 | B_13 |
| MOTA | 1778 | N | LYS | 33 | 66.021 | 68.461 | 10.636 | 1.00 12.52 | |
| MOTA | 1780 | | LYS | 33 | 65.663 | 69.786 | 10.171 | | B_13 |
| ATOM | 1781 | | LYS | 33 | 66.889 | 70.592 | | 1.00 13.00 | B_13 |
| ATOM | 1782 | | LYS | 33 | | | 9.762 | 1.00 22.63 | B_13 |
| ATOM | 1783 | | | | 66.581 | 72.054 | 9.560 | 1.00 18.24 | B_13 |
| | | | LYS | 33 | 65.604 | 72.545 | 10.630 | 1.00 29.21 | B_13 |
| MOTA | 1784 | | LYS | 33 | 66.185 | 72.429 | 12.048 | 1.00 41.79 | B_13 |
| MOTA | 1785 | | LYS | 33 | 65.181 | 71.939 | 13.054 | 1.00 20.17 | B_13 |
| MOTA | 1789 | | LYS | 33 | 64.698 | 69.686 | 9.023 | 1.00 10.62 | B_13 |
| MOTA | 1790 | | LYS | 33 | 63.734 | 70.437 | 8.971 | 1.00 22.94 | B_13 |
| MOTA | 1791 | N | ALA | 34 | 64.915 | 68.707 | 8.150 | 1.00 10.00 | B_13 |
| MOTA | 1793 | CA | ALA | 34 | 64.050 | 68.475 | 7.000 | 1.00 11.94 | B_13 |
| MOTA | 1794 | CB | ALA | 34 | 64.611 | 67.374 | 6.100 | 1.00 10.00 | B_13 |
| ATOM | 1795 | C | ALA | 34 | 62.640 | 68.115 | 7.423 | 1.00 10.00 | B_13 |
| ATOM | 1796 | | ALA | 34 | 61.675 | 68.650 | 6.878 | 1.00 15.32 | |
| MOTA | 1797 | | PHE | 35 | 62.510 | | | | B_13 |
| ATOM | 1799 | | PHE | 35 | | 67.208 | 8.387 | 1.00 21.32 | B_13 |
| ATOM | 1800 | | | | 61.187 | 66.789 | 8.852 | 1.00 18.32 | B_13 |
| | | | PHE | 35 | 61.267 | 65.451 | 9.614 | 1.00 25.48 | B_13 |
| MOTA | 1801 | | PHE | 35 | 61.620 | 64.260 | 8.735 | 1.00 14.33 | B_13 |
| ATOM | 1802 | | PHE | 35 . | 61.149 | 64.171 | 7.427 | 1.00 17.91 | B_13 |
| ATOM | 1803 | CD2 | | 35 | 62.436 | 63.240 | 9.217 | 1.00 18.05 | B_13 |
| ATOM | 1804 | CE1 | PHE | 35 | 61.486 | 63.086 | 6.610 | 1.00 18.49 | B_13 |
| MOTA | 1805 | CE2 | | 35 | 62.778 | 62.158 | 8.413 | 1.00 15.01 | |
| ATOM | 1806 | | PHE | 35 | 62.301 | 62.081 | | | B_13 |
| ATOM | 1807 | | PHE | 35 | 60.428 | | 7.103 | 1.00 10.00 | B_13 |
| ATOM | 1808 | | PHE | | | 67.862 | 9.658 | 1.00 18.68 | B_13 |
| ATOM | 1809 | | | 35 | 59.202 | 67.971 | 9.556 | 1.00 17.05 | B_13 |
| | | | LYS | 36 | 61.160 | 68.664 | 10.425 | 1.00 16.30 | B_13 |
| ATOM | 1811 | | LYS | 36 | 60.579 | 69.749 | 11.229 | 1.00 19.34 | B_13 |
| MOTA | 1812 | | LYS | 36 | 61.676 | 70.420 | 12.052 | 1.00 24.61 | B_13 |
| MOTA | 1813 | | LYS | 36 | 61.200 | 71.293 | 13.191 | 1.00 18.38 | B_13 |
| ATOM | | 77 | LYS | 36 | £3 400 | 71 705 | | | |
| | 1814 | | | | 62.408 | 71.795 | 13.962 | 1.00 19.34 | B 13 |
| MOTA | 1814 | | LYS | 36 | 62.408 | 72.267 | 15.356 | 1.00 19.34 | B_13 B_13 |

| 1001 | 1016 | | | 2.6 | | | | | • |
|-------|------|------|-----|------------|--------|----------|--------|-------------|-------------|
| ATOM | 1816 | NZ | LYS | 36 | 63.299 | 72.615 | 16.118 | 1.00 27.76 | B_13 |
| ATOM | 1320 | С | LYS | 36 | 59.924 | 70.770 | 10.301 | 1.00 10.19 | B_13 |
| MOTA | 1821 | 0 | LYS | 36 | 58.788 | 71.183 | 10.528 | 1.00 14.95 | B_13 |
| MOTA | 1822 | N | LYS | 37 | 60.630 | 71.134 | 9.233 | 1.00 15.89 | B_13 |
| ATOM | 1824 | .CA | LYS | 37 | 60.126 | 72.076 | 8.230 | 1.00 19.95 | B_13 |
| ATOM | 1825 | CB | LYS | 37 | 61.202 | 72.386 | 7.189 | 1.00 10.00 | |
| ATOM | 1826 | CG | LYS | 37 | | | | | B_13 |
| | | | | | 62.209 | 73.439 | 7.569 | 1.00 13.18 | B_13 |
| ATOM | 1827 | CD | LYS | 37 | 62.869 | 73.966 | 6.311 | 1.00 28.86 | B_13 |
| ATOM | 1828 | CE | LYS | 37 | 61.825 | 74.460 | 5.281 | 1.00 31.44 | B_13 |
| MOTA | 1829 | NZ | LYS | 37 | 60.878 | 75.512 | 5.772 | 1.00 26.23 | B_13 |
| MOTA | 1833 | C | LYS | 37 | 58.939 | 71.482 | 7.472 | 1.00 25.64 | |
| ATOM | 1834 | Ö | LYS | 37 | | | | | B_13 |
| | | | | | 57.968 | 72.177 | 7.161 | 1.00 24.39 | B_13 |
| ATOM | 1835 | - N | ALA | 38 | 59.060 | 70.205 | 7.128 | 1.00 17.12 | B_13 |
| MOTA | 1837 | CA | ALA | 38 | 58.031 | 69.493 | 6.381 | 1.00 16.06 | B_13 |
| MOTA | 1838 | CB | ALA | 38 | 58.459 | 68.038 | 6.154 | 1.00 12.19 | B_13 |
| ATOM | 1839 | С | ALA | 38 | 56.692 | 69.557 | 7.094 | 1.00 11.12 | B_13 |
| MOTA | 1840 | 0 | ALA | 38 | 55.648 | 69.736 | 6.458 | 1.00 31.10 | |
| MOTA | 1841 | Ň | PHE | 39 | | | | | B_13 |
| | | | | | 56.732 | 69.393 | 8.417 | 1.00 21.01 | B_13 |
| MOTA | 1843 | CA | PHE | 39 | 55.540 | 69.446 | 9.257 | 1.00 10.85 | B_13 |
| ATOM | 1844 | CB | PHE | 39 | 55.841 | 68.833 | 10.639 | 1.00 14.45 | B_13 |
| ATOM | 1845 | ::CG | PHE | 39 | 55.851 | 67.325 | 10.659 | 1.00 21.88 | B_13 |
| ATOM | 1846 | CD1 | PHE | 39 | 57.016 | 66.625 | 10.954 | 1.00 16.88 | B_13 |
| MOTA | 1847 | CD2 | | 39 | 54.675 | 66.599 | | | |
| ATOM | 1848 | CE1 | | | | | 10.442 | 1.00 22.14 | B_13 |
| | | | | 3 <i>9</i> | 57.010 | 65.223 | 11.037 | 1.00 17.95 | B_13 |
| MOTA | 1849 | CE2 | | 39 | 54.655 | 65.190 | 10.522 | 1.00 17.22 | B_13 |
| MOTA | 1850 | CZ | PHE | 39 | 55.823 | 64.503 | 10_823 | 1.00 13.51 | B_13 |
| ATOM. | 1851 | C | PHE | 39 | 55.044 | 70.898 | 9.426 | 1.00 19.98 | B_13 |
| MOTA | 1852 | Ö | PHE | 39 | 53.839 | 71.160 | 9.393 | 1.00 14.30 | _ |
| ATOM | 1853 | N | LYS | 40 | 55.981 | | | | B_13 |
| | | | | | | 71.826 | 9.611 | 1.00 20.03 | B_13 |
| MOTA | 1855 | CA | LYS | 40 | 55.681 | 73.245 | 9.795 | 1.00 18.64 | B_13 |
| MOTA | 1856 | CB | LYS | 40 | 56.989 | 74.011 | 10.020 | 1.00 19.28 | B_13 |
| ATOM | 1857 | CG | LYS | 40 | 57.064 | 75.392 | 9.440 | 1.00 26.34 | B_13 |
| MOTA | 1858 | CD | LYS | 40 | 58.288 | 76.093 | 9.974 | 1.00 18.46 | _ |
| MOTA | 1859 | CE | LYS | 40 | 58.021 | 76.673 | 11.339 | | B_13 |
| MOTA | 1860 | NZ | | | | | | 1.00 20.86 | B_13 |
| | | | LYS | 40 | 57.053 | 77.814 | 11.232 | 1.00 27.28 | B_13 |
| MOTA | 1864 | С | LYS | 40 | 54.899 | 73.790 | 8.612 | 1.00 20.57 | B_13 |
| MOTA | 1865 | 0 | LYS | 40 | 54.034 | 74.654 | 8.756 | 1.00 22.54 | B_13 |
| ATOM | 1866 | N | VAL | 41 | 55.216 | 73.251 | 7.445 | 1.00 17.15 | B_13 |
| MOTA | 1868 | CA | VAL | 41 | 54.565 | 73.576 | | | |
| ATOM | 1869 | CB | | | | | 6.184 | 1.00 19.19 | B_13 |
| | | | VAL | 41 | 55.095 | 72.566 | 5.086 | 1.00 17.28 | B_13 |
| MOTA | 1870 | | VAL | 41 | 53.987 | 72.064 | 4.160 | 1.00 10.00 | B_13 |
| MOTA | 1871 | CG2 | VAL | 41 | 56.224 | 73.191 | 4.293 | 1.00 19.38 | · B_13 |
| MOTA | 1872 | C | VAL | 41 | 53.026 | 73.472 | 6.354 | 1.00 20.38 | B_13 |
| MOTA | 1873 | 0 | VAL | 41 | 52.268 | 74.280 | 5.810 | 1.00 28.57 | |
| MOTA | 1874 | N | TRP | 42 | | | | | B_13 |
| | | | | | 52.587 | 72.511 | 7.163 | 1.00 23.10 | B_13 |
| ATOM | 1876 | CA | TRP | 42 | 51.166 | 72.265 | 7.403 | 1.00 19.29 | B_13 |
| MOTA | 1877 | CB | TRP | 42 | 50.912 | 70.757 | 7.487 | 1.00 22.19 | , B_13 |
| MOTA | 1878 | CG | TRP | 42 | 51.437 | 70.007 | 6.313 | 1.00 19.32 | B_13 |
| MOTA | 1879 | CD2 | TRP | 42 | 50.836 | 69.909 | 5.015 | 1.00 31.02 | B_13 |
| MOTA | 1880 | CE2 | TRP | 42 | 51.659 | 69.067 | 4.238 | 1.00 22.49. | |
| MOTA | 1881 | CE3 | | 42 | 49.677 | 70.448 | | | |
| ATOM | 1882 | CD1 | | | | | 4.434 | 1.00 15.54 | B_13 |
| | | | | 42 | 52.571 | 69.251 | 6.269 | 1.00 14.04 | B_13 |
| MOTA | 1883 | NE1 | | 42 | 52.710 | 68.681 | 5.027 | 1.00 13.55 | B_13 |
| ATOM | 1885 | CZ2 | | 42 | 51.360 | . 68.752 | 2.912 | 1.00 18.87 | B_13 |
| MOTA | 1886 | CZ3 | | 42 | 49.383 | 70.132 | 3.116 | 1.00 13.33 | B_13 |
| MOTA | 1887 | CH2 | TRP | 42 | 50.219 | 69.294 | 2.370 | 1.00 20.30 | B_13 |
| ATOM | 1888 | C | TRP | 42 | 50.617 | 72.926 | 8.660 | 1.00 24.68 | |
| MOTA | 1889 | Ō | TRP | 42 | 49.455 | 73.339 | | | B_13 |
| MOTA | 1890 | N | | | | | 8.688 | 1.00 20.93 | B_13 |
| | | | SER | 43 | 51.432 | 72.987 | 9.710 | 1.00 20.63 | B_13 |
| MOTA | 1892 | CA | SER | 43 | 51.007 | 73.601 | 10.968 | 1.00 22.47 | B_13 |
| MOTA | 1893 | CB | SER | 43 | 51.955 | 73.231 | 12.116 | 1.00 10.00 | B_13 |
| MOTA | 1894 | OĢ | SER | 43 | 53.265 | 73.716 | 11.891 | 1.00 33.50 | B_13 |
| MOTA | 1896 | C | SER | 43 | 50.913 | 75.122 | 10.829 | 1.00 14.99 | |
| ATOM | 1897 | 0 | SER | 43 | 50.224 | | | | B_13 |
| ATOM | 1898 | N | | | | | 11.595 | 1.00 11.58 | B_13 |
| | | | ASP | 44 | 51.613 | 75.667 | 9.843 | 1.00 26.20 | B_13 |
| ATOM | 1900 | CA | ASP | 44 | 51.595 | 77.100 | 9.617 | 1.00 22.11 | B_13 |
| ATOM | 1901 | CB | ASP | 44 | 52.620 | 77.485 | 8.549 | 1.00 11.09 | B_13 |
| ATOM | 1902 | CG | ASP | 44 | 54.000 | 77.751 | 9.125 | 1.00 18.45 | B_13 |
| MOTA | 1903 | | ASP | 44 | 54.903 | 78.114 | 8.347 | 1.00 17.67 | |
| MOTA | 1904 | | ASP | 44 | | | | · | B_13 |
| MOTA | 1905 | | | | 54.195 | 77.602 | 10.345 | 1.00 21.44 | B_13 |
| | | C | ASP | 44 | 50.216 | 77.575 | 9.190 | 1.00 32.83 | B_13 |
| MOTA | 1906 | 0 | ASP | 44 | 49.795 | 78.677 | 9.549 | 1.00 34.78 | B_13 |
| MOTA | 1907 | N | VAL | 45 | 49.508 | 76.735 | 8.439 | 1.00 31.40 | B_13 |
| MOTA | 1909 | CA | VAL | 45 | 48.191 | 77.094 | 7.932 | 1.00 14.00 | B_13 |
| MOTA | 1910 | CB | VAL | 45 | 48.121 | 76.872 | 6.401 | 1.00 15.73 | |
| MOTA | 1911 | | VAL | 45 | 49.123 | 77.755 | 5.707 | | B_13 |
| ATOM | 1912 | | VAL | 45 | | | | 1.00 19.37 | B_13 |
| | | -32 | 444 | 70 | 48.407 | 75.409 | 6.055 | 1.00 10.00 | B_13 |
| | | | | | | | | | |

| ATOM | 1913. | С | VAL | 45 | 47.054 | 76.333 | 8.575 | 1.00 18.43 | D 13 |
|------|-------------------|-----|-----|-----|----------|--------|--------|------------|-------------|
| ATCM | 1914 | Ō | VAL | 45 | | | | | B_13 |
| | | | | | 45.954 | 76.304 | 8.026 | 1.00 26.09 | B_13 |
| MOTA | 1915 | N | THR | 46 | 47.295 | 75.754 | 9.747 | 1.00 18.49 | B_13 |
| MOTA | 1917 | CA | THR | 46 | 46.262 | 74.963 | 10.408 | 1.00 21.92 | B_13 |
| ATOM | 1918 | CB | THR | 46 | | | | | |
| | | | | | 46.222 | 73.529 | 9.751 | 1.00 27.61 | B_13 |
| ATOM | 1919 | OG1 | THR | 46 | 44.876 | 73.047 | 9.661 | 1.00 28.78 | B_13 |
| MOTA | 1921 | CG2 | THR | 46 | 47.054 | 72.550 | 10.522 | 1.00 10.65 | B_13 |
| ATOM | 1922 | C | THR | 46 | | | | | |
| | | | | | 46.505 | 74.931 | 11.932 | 1.00 18.41 | B_13 |
| ATOM | 1923 | 0 | THR | 46 | 47.554 | 75.363 | 12.411 | 1.00 18.63 | B_13 |
| MOTA | 1924 | N | PRO | 47 | 45.519 | 74.467 | 12.717 | 1.00 16.81 | _ |
| | | | | | | | | | B_13 |
| ATOM | 1925 | CD | PRO | 47 | 44.113 | 74.209 | 12.348 | 1.00 32.80 | B_13 |
| MOTA | 1926 | CA | PRO | 47 | 45.691 | 74.407 | 14.169 | 1.00 13.66 | B_13 |
| MOTA | 1927 | CB | PRO | 47 | 44.256 | 74.489 | 14.675 | 1.00 30.52 | |
| | | | | | | | | | B_13 |
| MOTA | 1928 | CG | PRO | 47 | 43.519 | 73.692 | 13.638 | 1.00 29.25 | B_13 |
| MOTA | 1929 | С | PRO | 47 | 46.346 | 73.105 | 14.622 | 1.00 28.40 | B_13 |
| ATOM | 1930 | 0 | PRO | 47 | 46.037 | 72.597 | | - | |
| | | _ | | | | | 15.705 | 1.00 29.19 | B_13 |
| ATOM | 1931 | N | LEU | 48 | 47.220 | 72.547 | 13.784 | 1.00 27.10 | B_13 |
| ATOM | 1933 | CA | LEU | 48 | 47.915 | 71.302 | 14.124 | 1.00 21.49 | B_13 |
| MOTA | 1934 | CB | LEU | 48 | 48.087 | | | | |
| | | | | | | 70.418 | 12.885 | 1.00 16.21 | B_13 |
| MOTA | 1935 | CG | LEU | 48 | 46.924 | 69.476 | 12.538 | 1.00 15.14 | B_13 |
| MOTA | 1936 | CD1 | LEU | 48 | 45.618 | 70.049 | 13.000 | 1.00 26.83 | B_13 |
| MOTA | 1937 | CD2 | LEU | 48 | 46.894 | 69.206 | 11.035 | 1.00 32.93 | |
| ATOM | | | | | | | | | B_13 |
| | 1938 | C | LEU | 48 | 49.262 | 71.611 | 14.771 | 1.00 16.35 | B_13 |
| MOTA | 1939 | 0 | LEU | 48 | 49.885 | 72.648 | 14.498 | 1.00 26.65 | B_13 |
| ATOM | 1940 | N | ASN | 49 | 49.691 | 70.744 | 15.669 | 1.00 18.84 | _ |
| | | | | | | | | | B_13 |
| MOTA | 1942 | CA | ASN | 49 | 50.956 | 70.940 | 16.354 | 1.00 25.67 | B_13 |
| MOTA | 1943 ⁻ | CB | ASN | 49 | 50.741 | 71.205 | 17.846 | 1.00 23.64 | B_13 |
| MOTA | 1944 | CG | ASN | 49 | 49.734 | 72.301 | 18.100 | 1.00 23.64 | |
| MOTA | | | | | | | | | B_13 |
| | 1945 | OD1 | ASN | 49 | 48.895 | 72.192 | 18.989 | 1.00 33.47 | B_13 |
| MOTA | 1946 | ND2 | ASN | 49 | 49.796 | 73.359 | 17.305 | 1.00 37.40 | B_13 |
| MOTA | 1949 | ,C | ASN | 49 | 51.695 | 69.643 | 16.195 | 1.00 22.08 | |
| | _ | - | | | | | | | B_13 |
| MOTA | 1950 | 0 | ASN | 49 | 51.087 | 68.577 | 16.252 | 1.00 23.48 | B_13 |
| ATOM | 1951 | N | PHE | 50 | 52.994 | 69.723 | 15.951 | 1.00 25.59 | B_13 |
| ATOM | 1953 | CA | PHE | 50 | 53.762 | 68.510 | 15.806 | 1.00 19.57 | _ |
| MOTA | 1954 | CB | | | | | | | B_13 |
| | | | PHE | 50 | 54.258 | 68.343 | 14.380 | 1.00 12.47 | B_13 |
| ATOM | 1955 | CG | PHE | 50 | 53.161 | 68.024 | 13.432 | 1.00 14.47 | B_13 |
| MOTA | 1956 | CD1 | PHE | 50 | 52.665 | 68.989 | 12.581 | 1.00 17.81 | |
| MOTA | 1957 | CD2 | PHE | | | | | | B_13 |
| | | | | 50 | 52.566 | 66.770 | 13.445 | 1.00 14.44 | ₽_13 |
| ATOM | 1958 | CE1 | PHE | 50 | 51.585 | 68.705 | 11.754 | 1.00 23.43 | B_13 |
| MOTA | 1959 | CE2 | PHE | 50 | 51.488 | 66.482 | 12.624 | 1.00 20.62 | |
| MOTA | 1960 | CZ | PHE | | | | | | B_13 |
| | | | | 50 | 50.999 | 67.447 | 11.781 | 1.00 13.34 | B_13 |
| ATOM | 1961 | C | PHE | 50 | . 54.858 | 68.419 | 16.826 | 1.00 23.56 | B_13 |
| MOTA | 1962 | 0 | PHE | 50 | 55.720 | 69.299 | 16.922 | 1.00 20.28 | |
| ATOM | 1963 | N | | | | | | | B_13 |
| | | | THR | 51 | 54.728 | 67.387 | 17.651 | 1.00 26.45 | B_13 |
| ATOM | 1965 | CA | THR | 51 | 55.650 | 67.090 | 18.725 | 1.00 29.37 | B_13 |
| MOTA | 1966 | CB | THR | 51 | 54.851 | 66.834 | 20.024 | 1.00 28.17 | B_13 |
| MOTA | 1967 | OG1 | THR | 51 | | | | | |
| | | | | | 53.946 | 65.738 | 19.824 | 1.00 40.86 | B_13 |
| MOTA | 1969 | CG2 | THR | 51 | 54.032 | 68.078 | 20.393 | 1.00 25.37 | B_13 |
| MOTA | 1970 | C | THR | 51 | 56.435 | 65.838 | 18.331 | 1.00 21.26 | B_13 |
| MOTA | 1971 | 0 | THR | 51 | 55.849 | | 17.882 | | |
| ATOM | 1972 | | | | | | | 1.00 17.45 | B_13 |
| | | N | ARG | 52 | 57.755 | 65.889 | 18.477 | 1.00 15.17 | B_13 |
| MOTA | 1974 | CA | ARG | 52 | 58.604 | 64.752 | 18.126 | 1.00 20.79 | B_13 |
| MOTA | 1975 | CB | ARG | 52 | 59.868 | 65.241 | 17.429 | 1.00 20.81 | |
| ATOM | 1976 | CG | ARG | | | | | | B_13 |
| | | | | 52 | 60.871 | 64.160 | 17.110 | 1.00 19.06 | B_13 |
| MOTA | 1977 | CD | ARG | 52 | 62.208 | 64.808 | 16.880 | 1.00 22.17 | B_13 |
| ATOM | 1978 | NE | ARG | 52 | 63.293 | 63.848 | 16.904 | 1.00 18.57 | B_13 |
| MOTA | 1980 | CZ | ARG | 52 | 64.563 | 64.160 | | | |
| | | | | | | | 17.108 | 1.00 10.00 | B_13 |
| MOTA | 1981 | | ARG | 52 | 64.915 | 65.414 | 17.315 | 1.00 19.35 | B_13 |
| MOTA | 1984 | NH2 | ARG | 52 | 65.488 | 63.214 | 17.039 | 1.00 35.90 | B_13 |
| ATOM | 1987 | С | ARG | 52 | 58.995 | 63.903 | 19.328 | 1.00 22.29 | |
| | | | | | | | | _ | B_13 |
| MOTA | 1988 | 0 | ARG | 52 | 59.326 | 64.433 | 20.387 | 1.00 24.98 | B_13 |
| MOTA | 1989 | N . | LEU | 53 | 59.013 | 62.586 | 19.140 | 1.00 19.90 | B_13 |
| MOTA | 1991 | CA | LEU | 53 | 59.378 | 61.660 | | | |
| | | | | | | | 20.203 | 1.00 27.02 | B_13 |
| ATOM | 1992 | CB | LEU | 53 | 58.279 | 60.625 | 20.434 | 1.00 16.80 | B_13 |
| MOTA | 1993 | CG | LEU | 53 | 56.859 | 61.138 | 20.639 | 1.00 23.45 | B_13 |
| MOTA | 1994 | | LEU | 53 | 55.943 | 59.943 | | _ | |
| MOTA | | | | | | | 20.884 | 1.00 24.07 | B_13 |
| | 1995 | | LEU | 53 | 56.801 | 62.143 | 21.785 | 1.00 21.02 | B_13 |
| MOTA | 1996 | C | LEU | 53 | 60.657 | 60.944 | 19.813 | 1.00 15.08 | B_13 |
| MOTA | 1997 | 0 | LEU | 53 | 60.822 | 60.539 | 18.671 | | |
| MOTA | | | | | | | | 1.00 13.89 | B_13 |
| | 1998 | N | HIS | 54 | 61.532 | 60.750 | 20.792 | 1.00 19.96 | B_13 |
| MOTA | 2000 | CA | HIS | 54 | 62.812 | 60.079 | 20.568 | 1.00 28.80 | B_13 |
| ATOM | 2001 | CB | HIS | 54 | 63.848 | 60.604 | 21.569 | 1.00 19.40 | |
| ATOM | | | | | | | | | B_13 |
| | 2002 | CG | HIS | 54 | 64.113 | 62.075 | 21.431 | 1.00 31.96 | B_13 |
| ATOM | 2003 | CD2 | HIS | 54 | 63.365 | 63.060 | 20.883 | 1.00 21.32 | B_13 |
| MOTA | 2004 | ND1 | HIS | 54 | 65.292 | 62.662 | 21.835 | 1.00 33.94 | B_13 |
| ATOM | 2006 | | HIS | | | | | _ | |
| | | | | .54 | 65.260 | 63.949 | 21.539 | 1.00 18.64 | B_13 |
| ATOM | 2007 | NE2 | HIS | 54 | 64.103 | 64.218 | 20.960 | 1.00 19.56 | B_13 |

| MOTA | 2009 | С | HIS | 54 | 62.695 | 58.555 | 20.647 | 1.00 13.04 | B_13 |
|--------|------|-----|-------|----------|--------|--------|--------|------------|------|
| ATOM | 2010 | 0 | HIS | 54 | 63.620 | 57.850 | 20.282 | 1.00 19.90 | B_13 |
| MOTA | 2011 | N | ASP | 55 | 61.586 | 58.076 | 21.219 | 1.00 17.27 | B_13 |
| MOTA | 2013 | CA | ASP | 55 | 61.303 | 56.648 | 21.366 | 1.00 25.79 | B_13 |
| MOTA | 2014 | CB | ASP | 55 | 62.099 | 56.038 | 22.533 | 1.00 29.40 | B_13 |
| MOTA | 2015 | CG | ASP | 55 | 63.443 | 55.428 | 22.076 | 1.00 29.64 | B_13 |
| MOTA | 2016 | OD1 | ASP | 55 | 63.517 | 54.906 | 20.942 | 1.00 33.28 | B_13 |
| ATOM | 2017 | | ASP | 55 | 64.437 | 55.469 | 22.831 | 1.00 31.99 | B_13 |
| ATOM | 2018 | C | ASP | 55 | 59.807 | 56.460 | 21.567 | 1.00 24.99 | B_13 |
| ATOM | 2019 | Õ | ASP | 55 | 59.079 | 57.445 | | | |
| | | _ | | | | | 21.677 | 1.00 21.06 | B_13 |
| MOTA | 2020 | N | GLY | 56 | 59.358 | 55.207 | 21.559 | 1.00 22.90 | B_13 |
| MOTA | 2022 | CA | GLY | 56 | 57.954 | 54.877 | 21.737 | 1.00 21.80 | B_13 |
| MOTA | 2023 | C | GLY | 56 | 57.155 | 54.926 | 20.447 | 1.00 14.48 | B_13 |
| MOTA | 2024 | 0 | GLY | 56 | 57.720 | 55.108 | 19.379 | 1.00 19.38 | B_13 |
| ATOM | 2025 | N | ILE | 57 | 55.841 | 54.742 | 20.545 | 1.00 11.78 | B_13 |
| MOTA | 2027 | CA | ILE | 57 | 54.944 | 54.809 | 19.389 | 1.00 16.25 | B_13 |
| MOTA | 2028 | CB | ILE | 57 | 53.737 | 53.804 | 19.510 | 1.00 22.94 | B_13 |
| MOTA | 2029 | CG2 | ILE | 57 | 52.442 | 54.417 | 18.955 | 1.00 24.79 | B_13 |
| MOTA | 2030 | CG1 | ILE | 57 | 54.025 | 52.505 | 18.744 | 1.00 25.63 | B_13 |
| MOTA | 2031 | CD1 | ILE | 57 | 53.586 | 52.520 | 17.240 | 1.00 17.48 | B_13 |
| ATOM | 2032 | C | ILE | 57 | 54.410 | 56.238 | 19.301 | 1.00 18.78 | B_13 |
| ATOM | 2033 | ŏ | ILE | 57 | 53.866 | 56.777 | 20.270 | 1.00 11.40 | |
| ATOM | 2034 | N | ALA | 58 | 54.598 | 56.842 | | | B_13 |
| ATOM | 2036 | CA | ALA | | | | 18.140 | 1.00 14.67 | B_13 |
| | | | | 58 50 | 54.139 | 58.200 | 17.857 | 1.00 17.04 | B_13 |
| ATOM | 2037 | CB | ALA | 58 | 55.270 | 59.015 | 17.245 | 1.00 10.00 | B_13 |
| MOTA | 2038 | C | ALA | 58 | 53.048 | 58.009 | 16.825 | 1.00 25.41 | B_13 |
| MOTA | 2039 | 0 | ALA | 58 | 52.956 | 56.940 | 16.243 | 1.00 22.59 | B_13 |
| MOTA | 2040 | N | ASP | 59 | 52.211 | 59.020 | 16.609 | 1.00 13.36 | B_13 |
| MOTA | 2042 | CA | ASP | 59 | 51.156 | 58.927 | 15.606 | 1.00 24.67 | B_13 |
| MOTA | 2043 | CB | ASP | 59 | 50.348 | 60.237 | 15.545 | 1.00 10.00 | B_13 |
| ATOM | 2044 | CG | ASP | 59 | 49.743 | 60.631 | 16.899 | 1.00 12.93 | B_13 |
| MOTA | 2045 | | ASP | 59 | 49.922 | 61.788 | 17.327 | 1.00 32.89 | B_13 |
| ATOM | 2046 | OD2 | | 59 | 49.076 | 59.793 | 17.541 | 1.00 21.52 | B_13 |
| MOTA | 2047 | C | ASP | 59 | 51.784 | 58.653 | 14.242 | 1.00 11.46 | B_13 |
| MOTA | 2048 | Ö | ASP | 59 | 51.378 | 57.736 | 13.531 | 1.00 16.58 | |
| ATOM | 2049 | N | ILE | 60 | | | | | B_13 |
| | | | | | 52.791 | 59.445 | 13.899 | 1.00 24.90 | B_13 |
| MOTA | 2051 | CA | ILE | 60 | 53.494 | 59.346 | 12.624 | 1.00 12.17 | B_13 |
| MOTA | 2052 | CB | ILE | 60 | 53.620 | 60.738 | 11.975 | 1.00 10.91 | B_13 |
| ATOM | 2053 | CG2 | | 60 | 54.289 | 60.641 | 10.588 | 1.00 10.70 | B_13 |
| ATOM | 2054 | CG1 | | 60 | 52.228 | 61.367 | 11.851 | 1.00 18.58 | B_13 |
| MOTA | 2055 | CD1 | ILE | 60 | 52.219 | 62.870 | 11.726 | 1.00 12.00 | B_13 |
| ATOM | 2056 | С | ILE | 60 | 54.881 | 58.750 | 12.841 | 1.00 12.93 | B_13 |
| MOTA | 2057 | 0 | ILE | 60 · | 55.788 | 59.392 | 13.365 | 1.00 16.39 | B_13 |
| ATOM | 2058 | N | MET | 61 | 55.015 | 57.485 | 12.483 | 1.00 19.08 | B_13 |
| MOTA | 2060 | CA | MET | 61 | 56.275 | 56.784 | 12.617 | 1.00 16.97 | B_13 |
| MOTA | 2061 | CB | MET | 61 | 56.011 | | 13.035 | 1.00 23.79 | B_13 |
| ATOM | 2062 | CG | MET | 61 | 55.313 | 55.172 | 14.422 | 1.00 12.37 | B_13 |
| MOTA | 2063 | SD | MET | 61 | 56.389 | 55.360 | 15.913 | 1.00 31.01 | |
| ATOM | 2064 | CE | MET | 61 | 57.204 | | | | B_13 |
| ATOM | 2065 | C | MET | 61 | _ | | | 1.00 14.93 | B_13 |
| ATOM | 2066 | | | | 56.995 | 56.888 | 11.265 | 1.00 12.72 | B_13 |
| | | 0 | MET | 61 | 56.438 | 56.538 | 10.216 | 1.00 15.31 | B_13 |
| ATOM | 2067 | N | ILE | 62 | 58.170 | 57.518 | 11.294 | 1.00 16.64 | B_13 |
| ATOM | 2069 | CA | ILE | 62 | 58.978 | 57.739 | | 1.00 27.48 | B_13 |
| ATOM | 2070 | CB | ILE. | - | 59.557 | 59.181 | 10.060 | 1.00 10.00 | B_13 |
| MOTA | 2071 | CG2 | | 62 | 60.191 | 59.462 | 8.717 | 1.00 18.65 | B_13 |
| ATOM | 2072 | CG1 | | 62 | 58.460 | 60.203 | 10.342 | 1.00 18.51 | B_13 |
| MOTA | 2073 | CD1 | | 62 | 58.983 | | 10.931 | 1.00 16.23 | B_13 |
| MOTA | 2074 | С | ILE | 62 | 60.155 | 56.787 | 10.046 | 1.00 15.06 | B_13 |
| MOTA | 2075 | 0 | ILE | 62· | 60.873 | 56.606 | 11.033 | 1.00 10.73 | B_13 |
| MOTA | 2076 | N | SER | 63 | 60.398 | 56.230 | 8.873 | 1.00 19.40 | B_13 |
| ATOM | 2078 | CA | SER | 63 | 61.513 | 55.321 | 8.722 | 1.00 13.31 | B_13 |
| MOTA | 2079 | CB | SER | 63 | 61.111 | | 9.123 | 1.00 17.28 | B_13 |
| MOTA | 2080 | OG | SER | 63 | 59.985 | | | 1.00 13.66 | B_13 |
| MOTA | 2082 | C | SER | 63 | 62.086 | | | 1.00 13.86 | |
| MOTA | 2083 | Õ | SER | 63 | | | | | B_13 |
| MOTA | 2083 | N | | | 61.441 | | | 1.00 20.93 | B_13 |
| | | | PHE | 64 | 63.338 | | | 1.00 17.78 | B_13 |
| ATOM | 2086 | CA | PHE | 64 | 64.072 | | | 1.00 18.81 | B_13 |
| ATOM | 2087 | CB | PHE | 64 | 65.409 | | 6.105 | 1.00 16.50 | B_13 |
| MOTA | 2088 | CG | PHE | 64 | 65.278 | _ | | 1.00 22.54 | B_13 |
| ATOM | 2089 | CD1 | | | 65.321 | | | 1.00 20.48 | B_13 |
| MOTA | 2090 | | PHE | | 65.155 | 57.708 | 7.395 | 1.00 24.76 | B_13 |
| MOTA | 2091 | | . PHE | | 65.246 | 59.207 | | | B_13 |
| MOTA | 2092 | CE2 | PHE | | 65.079 | | | 1.00 14.29 | B_13 |
| MOTA | 2093 | CZ | PHE | | 65.128 | | | 1.00 10.16 | B_13 |
| ATOM ' | 2094 | C | PHE | | 64.293 | | | 1.00 10.30 | B_13 |
| MOTA | 2095 | 0 | PHE | | 64.571 | | | | B_13 |
| MOTA | 2096 | N | GLY | 65 | 64.121 | | | | B_13 |
| - | | | | | | 12 | 010 | xJ.J0 | n_13 |

| » moss | 2000 | • | | | . | | | | |
|----------------|--------------|----------|------------|----------|------------------|------------------|-----------------|--------------------------|--------------|
| ATOM | 2098 | CA | GLY | 65 | 64.306 | 51.426 | 4.392 | 1.00 14.88 | B_13 |
| ATOM ATOM | 2099 2100 | C | GLY | 65 65 | 64.400 | 51.117 | 2.922 | 1.00 14.95 | B_13 |
| ATOM | 2100 | И | ILE | 65 66 | 64.047 | 51.947 | 2.088 | 1.00 12.61 | B_13 |
| ATOM | 2103 | CA | ILE | 66 | 64.860 | 49.922 | 2.587 | 1.00 10.00 | B_13 |
| ATOM | 2104 | CB | ILE | 66 | 64.995 66.483 | 49.555 49.344 | 1.187 | 1.00 19.70 1.00 18.92 | B_13 |
| ATOM | 2105 | CG2 | ILE | 66 | 67.301 | 50.628 | 0.791 1.073 | 1.00 18.92 | B_13 |
| MOTA | 2106 | CG1 | ILE | 66 | 67.078 | 48.178 | 1.582 | 1.00 10.00 | B_13 |
| MOTA | 2107 | CD1 | ILE | 66 | 68.381 | 47.662 | 1.004 | 1.00 14.64 | B_13 |
| MOTA | 2108 | C | ILE | 66 | 64.195 | 48.296 | 0.900 | 1.00 17.33 | B_13 |
| ATOM | 2109 | Õ | ILE | 66 | 63.877 | 47.543 | 1.806 | 1.00 20.10 | B_13 B_13 |
| ATOM | 2110 | N | LYS | 67 | 63.773 | 48.148 | -0.349 | 1.00 18.78 | B_13 |
| MOTA | 2112 | CA | LYS | 67 | 63.019 | 46.980 | -0.787 | 1.00 14.73 | B_13 |
| MOTA | 2113 | CB | LYS | 67 | 63.986 | 45.827 | -1.073 | 1.00 22.08 | B_13 |
| MOTA | 2114 | CG | LYS | 67 | 65.107 | 46.142 | -2.066 | 1.00 15.53 | B_13 |
| MOTA | 2115 | CD | LYS | 67 | 64.591 | 46.325 | -3.487 | 1.00 16.76 | B_13 |
| MOTA | 2116 | CE | LYS | 67 | 65.573 | 45.763 | -4.523 | 1.00 21.90 | B_13 |
| ATOM | 2117 | NZ | LYS | 67 | 66.975 | 46.257 | -4.394 | 1.00 28.03 | B_13 |
| ATOM | 2121 | C | LYS | 67 | 61.945 | 46.548 | 0.218 | 1.00 16.24 | B_13 |
| ATOM | 2122 | 0 | LYS | 67 | 61.136 | 47.360 | 0.649 | 1.00 10.25 | B_13 |
| ATOM | 2123 | N. | GLU | 68 | 61.968 | 45.293 | 0.630 | 1.00 10.00 | B_13 |
| MOTA ATOM | 2125 2126 | CA CB | GLU | 68 69 | 60.986 | 44.787 | 1.570 | 1.00 10.00 | B_13 |
| ATOM | 2127 | CG | GLU | 68 68 | 61.004 59.733 | 43.257 | 1.505 | 1.00 31.44 | B_13 |
| ATOM | 2128 | CD | GLU | 68 | 58.723 | 42.550 42.720 | 1.696 | 1.00 27.13 | B_13 |
| ATOM | 2129 | OE1 | GLU | 68 | 59.106 | 42.180 | 0.524 -0.613 | 1.00 12.88 1.00 14.05 | B_13 |
| ATOM | 2130 | OE2 | GLU | 68 | 57.681 | 43.274 | 0.753 | 1.00 14.05 | B_13 |
| ATOM | 2131 | C | GLU | 68 | 61.402 | 45.292 | 2.954 | 1.00 32.89 | B_13 B_13 |
| MOTA | 2132 | 0 | GLU | 68 | 62.541 | 45.099 | 3.390 | 1.00 19.77 | B_13 |
| MOTA | 2133 | N | HIS | 69 | 60.467 | 45.918 | 3.659 | 1.00 15.43 | B_13 |
| MOTA | 2135 | CA | HIS | 69 | 60.777 | 46.473 | 4.964 | 1.00 10.00 | B_13 |
| MOTA | 2136 | CB | HIS | 69 | 61.173 | 47.928 | 4.802 | 1.00 15.60 | B_13 |
| MOTA | 2137 | CG | HIS | 69 | 60.151 | 48.731 | 4.063 | 1.00 18.06 | B_13 |
| MOTA | 2138 | | HIS | 69 | 59.131 | 49.509 | 4.498 | 1.00 25.01 | B_13 |
| ATOM | 2139 | | HIS | 69 | 60.055 | 48.709 | 2.689 | 1.00 21.79 | B_13 |
| ATOM | 2141 | | HIS | 69 | 59.023 | 49.430 | 2.308 | 1.00 19.43 | B_13 |
| ATOM ATOM | 2142 | NEZ C | HIS | 69 60 | 58.438 | 49.932 | 3.384 | 1.00 19.23 | B_13 |
| ATOM | 2143 2144 | 0 | HIS HIS | 69 69 | 59.655 | 46.396 | 5.978 | 1.00 16.27 | B_13 |
| MOTA | 2145 | N | GLY | 70 | 59.689 58.610 | 47.099 45.629 | 6.969 5.719 | 1.00 13.47 | B_13 |
| ATOM | 2147 | CA | GLY | 70 | 57.567 | 45.520 | 6.720 | 1.00 21.21 1.00 15.93 | B_13 |
| ATOM | 2148 | C | GLY | 70 | 56.147 | 45.784 | 6.287 | 1.00 13.13 | B_13 B_13 |
| MOTA | 2149 | 0 | GLY | 70 | 55.283 | 45.986 | 7.147 | 1.00 12.19 | B_13 |
| ATOM | 2150 | N | ASP | 71 | 55.891 | 45.805 | 4.983 | 1.00 10.00 | B_13 |
| MOTA | 2152 | CA | ASP | 71 | 54.540 | 46.030 | 4.480 | 1.00 17.84 | B_13 |
| MOTA | 2153 | CB | ASP | 71 | 54.086 | 47.490 | 4.636 | 1.00 21.86 | B_13 |
| MOTA | 2154 | CG | ASP | 71 | 54.946 | 48.480 | 3.881 | 1.00 13.38 | B_13 |
| ATOM ATOM | 2155 | | ASP | 71 | 54.896 | 49.644 | 4.291 | 1.00 10.00 | B_13 |
| ATOM | 2156 2157 | C C | ASP ASP | 71 71 | 55.633 | 48.135 | 2.897 | 1.00 10.00 | B_13 |
| ATOM | 2158 | Ö | ASP | 71 | 54.313 55.221 | 45.557 45.068 | 3.064 | 1.00 27.18 | B_13 |
| ATOM | 2159 | N | PHE | 72 | 53.221 | 45.759 | 2.416 | 1.00 16.61 1.00 10.00 | B_13 |
| ATOM | 2161 | ÇA | PHE | 72 | 52.788 | 45.317 | 2.564 1.213 | 1.00 10.00 | B_13 B_13 |
| MOTA | 2162 | CB | PHE | 72 | 51.292 | 45.017 | 1.099 | 1.00 16.43 | B_13 |
| MOTA | 2163 | CG | PHE | 72 | 50.849 | 43.779 | 1.851 | 1.00 27.69 | B_13 |
| MOTA | 2164 | CD1 | PHE | 72 | 51.399 | 42.532 | 1.561 | 1.00 22.33 | B_13 |
| ATOM | 2165 | | PHE | 72 | 49.848 | 43.855 | 2.823 | 1.00 27.58 | B_13 |
| ATOM | 2166 | | PHE | 72 | 50.955 | 41.383 | 2.225 | 1.00 22.03 | B_13 |
| MOTA | 2167 | CE2 | | 72 | 49.403 | 42.709 | 3.486 | 1.00 21.82 | B_13 |
| ATOM | 2168 | CZ | PHE | 72 | 49.957 | 41.473 | 3.184 | 1.00 10.00 | B_13 |
| MOTA | 2169 | C | PHE | 72 72 | 53.225 | 46.313 | 0.130 | 1.00 18.56 | B_13 |
| MOTA MOTA | 2170 2171 | 0 | PHE | 72 73 | 52.840 | 46.190 | -1.048 | 1.00 14.78 | B_13 |
| ATOM | 2173 | N CA | TYR TYR | 73 73 | 54.079 | 47.260 | 0.513 | 1.00 10.93 | B_13 |
| MOTA | 2174 | CB | TYR | 73 73 | 54.558 53.943 | 48.295 | -0.416 | 1.00 13.87 | B_13 |
| MOTA | 2175 | CG | TYR | 73 | 52.439 | 49.649 49.581 | -0.048 | 1.00 22.69 | B_13 |
| ATOM | 2176 | | TYR | 73 | 51.774 | 49.385 | 0.007 1.219 | 1.00 16.43 1.00 18.21 | B_13 |
| ATOM | 2177 | | TYR | 73 | 50.386 | 49.219 | 1.219 | 1.00 18.21 | B_13 B_13 |
| MOTA | 2178 | | TYR | 73 | 51.683 | 49.618 | -1.165 | 1.00 35.13 | B_13 B_13 |
| MOTA | 2179 | CE2 | | 73 | 50.300 | 49.456 | -1.133 | 1.00 39.16 | B_13 |
| MOTA | 2180 | CZ | TYR | 73 | 49.663 | 49.258 | 0.080 | 1.00 28.27 | B_13 |
| MOTA | 2181 | OH | TYR | 73 | 48.301 | 49.122 | 0.106 | 1.00 33.06 | B_13 |
| ATOM | 2183 | C | TYR. | | 56.088 | 48.349 | -0.425 | 1.00 18.05 | B_13 |
| MOTA | 2184 | 0 | TYR | 73 | 56.721 | 49.339 | 0.003 | 1.00 10.00 | B_13 |
| MOTA MOTA | 2185 2186 | N | PRO | 74 | 56.702 | 47.287 | -0.953 | 1.00 13.76 | B_13 |
| MOTA | 2186 | CD CA | PRO PRO | 74 74 | 56.063 58.158 | 46.221 47.183 | -1.740 | 1.00 14.21 | B_13 |
| - ~ ~** | | ₩. | - 110 | | JU. 1J0 | 41.463 | -1.024 | 1.00 21.66 | B_13 |
| | | | | | | | | | |

| ATOM | 2188 | СВ | PRO | 74 | E0 3E3 | 45 760 | 1 560 | 1 00 15 00 | n 13 |
|--------------|--------------|-----------|------------|----------|------------------|------------------|-------------------|--------------------------|----------------|
| MOTA | 2189 | CG | PRO | 74 | 58.353 57.225 | 45.768 45.653 | -1.569 -2.540 | 1.00 15.88 1.00 13.95 | B_13 B_13 |
| ATOM | 2190 | C | PRO | 74 | 58.747 | 48.226 | -1.959 | 1.00 27.68 | B_13 |
| ATOM | 2191 | 0 | PRO | 74 | 58.173 | 48.526 | -3.012 | 1.00 21.90 | B_13 |
| ATOM | 2192 | N | PHE | 75 | 59.883 | 48.794 | -1.562 | 1.00 20.91 | B_13 |
| ATOM | 2194 | CA | PHE | 75 75 | 60.554 | 49.773 | -2.395 | 1.00 15.84 | B_13 |
| MOTA MOTA | 2195 2196 | CB CG | PHE | 75 75 | 61.498 60.765 | 50.637 | -1.548 | 1.00 11.67 | B_13 |
| ATOM | 2197 | CD1 | PHE | 75 75 | 59.831 | 51.589 52.484 | -0.641 -1.162 | 1.00 14.42 1.00 16.56 | B_13 B_13 |
| MOTA | 2198 | CD2 | PHE | 75 | 60.976 | 51.574 | 0.726 | 1.00 10.00 | B_13 |
| MOTA | 2199 | CE1 | PHE | 75 | 59.119 | 53.345 | -0.327 | 1.00 11.14 | B_13 |
| MOTA | 2200 | CE2 | PHE | 75 | 60.274 | 52.423 | 1.558 | 1.00 10.28 | B_13 |
| MOTA | 2201 | CZ | PHE | 75 | 59.340 | 53.316 | 1.027 | 1.00 10.00 | B_13 |
| ATOM | 2202 2203 | C | PHE | 75 75 | 61.236 | 49.068 | -3.573 | 1.00 14.23 | B_13 |
| MOTA MOTA | 2203 | N | PHE ASP | 75 76 | 61.357 61.742 | 47.837 49.845 | -3.582 -4.526 | 1.00 18.64 1.00 12.83 | B_13 · B_13 |
| ATOM | 2206 | CA | ASP | 76 | 62.330 | 49.287 | -5.740 | 1.00 20.69 | B_13 |
| MOTA | 2207 | CB | ASP | 76 | 61.394 | 49.644 | -6.911 | 1.00 14.28 | B_13 |
| MOTA | 2208 | CG | ASP | 76 | 61.212 | 51.144 | -7.080 | 1.00 14.37 | B_13 |
| MOTA | 2209 | OD1 | ASP | 76 26 | 61.361 | 51.882 | -6.095 | 1.00 22.32 | B_13 |
| MOTA MOTA | 2210 2211 | OD2 C | ASP ASP | 76 76 | 60.941 63.764 | 51.597 | -8.202 | 1.00 15.92 | B_13 |
| ATOM | 2212 | 0 | ASP | 76 | 64.056 | 49.698 49.864 | -6.104 -7.278 | 1.00 19.31 1.00 18.67 | B_13 B_13 |
| ATOM | 2213 | N. | GLY | 77 | 64.653 | 49.902 | -5.132 | 1.00 10.00 | B_13 |
| MOTA | 2215 | CA | GLY | 77 | 65.997 | 50.326 | -5.501 | 1.00 10.00 | B_13 |
| MOTA | 2216 | С | GLY | 77 | 65.989 | 51.790 | -5.970 | 1.00 16.22 | B_13 |
| MOTA | 2217 | 0 | GLY | 77 | 64.967 | 52.487 | -5.752 | 1.00 17.04 | B_13 |
| MOTA MOTA | 2218 2219 | N | PRO | 78 78 | 67.080 | 52.305 | -6.589 | 1.00 12.53 | B_13 |
| ATOM | 2220 | CD CA | PRO PRO | 78 78 | 68.319 67.207 | 51.564 53.691 | -6.856 -7.086 | 1.00 12.24 1.00 11.81 | B_13 B_13 |
| MOTA | 2221 | CB | PRO | 78 | 68.546 | 53.678 | -7.816 | 1.00 10.00 | B_13 |
| MOTA | 2222 | CG | PRO | 78 | 69.316 | 52.693 | -7.066 | 1.00 12.78 | B_13 |
| MOTA | 2223 | C | PRO | 78 | 66.093 | 54.146 | -8.027 | 1.00 10.00 | B_13 |
| MOTA | 2224 | 0 | PRO | 78 | 65.621 | 53.381 | -8.853 | 1.00 27.46 | B_13 |
| MOTA | 2225 | N | SER | 79 70 | 65.641 | 55.386 | -7.852 | 1.00 19.14 | B_13 |
| MOTA MOTA | 2227 2228 | CA CB | SER SER | 79 79 | 64.568 64.970 | 55.963 | -8.669 -10.148 | 1.00 10.00 1.00 20.11 | B_13 |
| ATOM | 2229 | OG | SER | 79 | 63.982 | | -10.148 | 1.00 23.87 | B_13 B_13 |
| ATOM | 2231 | C | SER | 79 | 63.231 | | -8.507 | 1.00 31.68 | B_13 |
| MOTA | 2232 | 0 | SER | 79 | 63.074 | 54.356 | -7.606 | 1.00 26.48 | B_13 |
| ATOM | 2233 | N | GLY | 80 | 62.250 | 55.589 | -9.327 | 1.00 10.00 | B_13 |
| MOTA | 2235 | CA | GLY | 80 | 60.940 | | -9.260 | 1.00 10.07 | B_13 |
| MOTA MOTA | 2236 2237 | C | GLY | 80 80 | 60.293 60.347 | 55.412 56.600 | -7.968 -7.643 | 1.00 30.72 1.00 20.65 | B_13 |
| ATOM | 2238 | N | LEU | 81 | 59.779 | | -7.193 | 1.00 20.05 | B_13 B_13 |
| ATOM | 2240 | CA | LEU | 81 | 59.135 | 54.752 | -5.917 | 1.00 13.14 | B <u>.</u> 13 |
| MOTA | 2241 | CB | LEU | 81 | 58.661 | 53.481 | -5.213 | 1.00 16.20 | B_13 |
| ATOM | 2242 | CG | LEU | 81 | 57.393 | 52.775 | -5.687 | 1.00 17.33 | B_13 |
| ATOM ATOM | 2243 2244 | | LEU | 81 | 57.554 | 52.277 | -7.096 | 1.00 28.67 | B_13 |
| ATOM | 2245 | CDZ | LEU | 81 81 | 57.103 60.122 | 51.617 55.466 | -4.745 -5.019 | 1.00 27.02 1.00 14.51 | B_13 B_13 |
| ATOM | 2246 | Õ | LEU | 81 | 61.264 | 55.016 | -4.846 | 1.00 16.24 | B_13 B_13 |
| MOTA | 2247 | N | LEU | 82 | 59.692 | 56.590 | -4.470 | 1.00 11.33 | B_13 |
| MOTA | 2249 | CA | LEU | 82 | 60.540 | 57.381 | -3.594 | 1.00 17.52 | B_13 |
| ATOM | 2250 | CB | LEU | 82 | 60.442 | 58.861 | -3.986 | 1.00 18.51 | B_13 |
| MOTA MOTA | 2251 2252 | CG CD1 | LEU LEU | 82 82 | 61.355 61.800 | 59.499 | -5.044 | 1.00 15.37 | B_13 |
| MOTA | 2253 | | LEU | 82 | 60.639 | 58.504 60.744 | -6.104 -5.659 | 1.00 17.05 1.00 16.87 | B_13 B_13. |
| ATOM | 2254 | C | LEU | 82 | 60.172 | 57.203 | -2.127 | 1.00 10.00 | B_13 |
| MOTA | 2255 | 0 | LEU | 82 | 61.045 | | -1.275 | 1.00 19.90 | B_13 |
| MOTA | 2256 | N | ALA | 83 | 58.876 | | -1.840 | 1.00 18.16 | B_13 |
| MOTA | 2258 | CA | ALA | 83 | 58.378 | | | 1.00 13.17 | B_13 |
| ATOM ATOM | 2259 2260 | CB | ALA | 83 | 58.762 | | | 1.00 10.00 | B_13 |
| MOTA | 2261 | 0 | ALA ALA | 83 83 | 56.846 56.209 | | | | B_13 |
| ATOM | 2262 | N | HIS | 84 | 56.268 | | | 1.00 10.73 | B_13 B_13 |
| MOTA | 2264 | CA | HIS | 84 | 54.811 | | | 1.00 23.81 | B_13 |
| MOTA | 2265 | CB | HIS | 84 | 54.270 | 55.188 | 0.157 | | B_13 |
| ATOM | 2266 | CG | HIS | 84 | 54.848 | | | 1.00 17.68 | B_13 |
| MOTA | 2267 | | HIS. | | 54.856 | | | 1.00 10.00 | B_13 |
| MOTA ATOM | 2268 2270 | | L HIS | 84 84 | 55.525 55.933 | | | | B_13 |
| MOTA | 2271 | | 2 HIS | 84 | 55.543 | | | 1.00 29.72 1.00 13.81 | B_13 B_13 |
| ATOM | 2272 | C | HIS | 84 | 54.363 | | | | B_13 |
| ATOM | 2273 | 0 | HIS | 84 | 55.099 | 56.148 | 3.166 | 1.00 20.02 | B_13 |
| MOTA | 2274 | N | ALA | 85 85 | 53.161 | | | | B_13 |
| MOTA | 2276 | CA | ALA | 85 | 52.584 | 57.230 | 3.796 | 1.00 18.64 | B_13 |

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| ATOM | 2277 | СВ | ALA | 85 | 52 620 | E0 705 | 4 222 | 1 00 13 00 | n 13 |
|--------------|--------------|----------|------------|----------------------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 2278 | C | ALA | 85 | 52.638 51.138 | 58.705 56.716 | 4.223 3.837 | 1.00 13.89 1.00 10.00 | B_13 |
| ATOM | 2279 | 0 | ALA | 85 | 50.434 | 56.728 | 2.828 | 1.00 10.00 | B_13 |
| ATOM | 2280 | N | PHE | 86 | 50.676 | 56.322 | 5.016 | 1.00 14.76 | B_13 B_13 |
| ATOM | 2282 | CA | PHE | 86 | 49.316 | 55.811 | 5.143 | 1.00 17.96 | B_13 |
| ATOM | 2283 | CB | PHE | 86 | 49.286 | 54.592 | 6.084 | 1.00 15.86 | B_13 |
| ATOM | 2284 | CG | PHE | 86 | 50.320 | 53.542 | 5.748 | 1.00 26.30 | B_13 |
| ATOM | 2285 | CD1 | PHE | 86 | 49.973 | 52.398 | 5.042 | 1.00 22.30 | B_13 |
| ATOM | 2286 | CD2 | PHE | 86 | 51.654 | 53.730 | 6.090 | 1.00 27.63 | B_13 |
| ATOM | 2287 | CE1 | PHE | 86 | 50.938 | 51.472 | 4.681 | 1.00 27.85 | B_13 |
| MOTA | 2288 | CE2 | PHE | 86 | 52.620 | 52.810 | 5.731 | 1.00 13.97 | B_13 |
| ATOM . | 2289 | CZ | PHE | 86 | 52.266 | 51.683 | 5.027 | 1.00 23.08 | B_13 |
| ATOM | 2290 | C | PHE | 86 | 48.427 | 56.924 | 5.669 | 1.00 13.02 | B_13 |
| ATOM | 2291 | Ö | PHE | 86 | 48.870 | 57.747 | 6.466 | 1.00 15.02 | B_13 |
| ATOM | 2292 | N | PRO | 87 | 47.174 | 57.006 | 5.186 | 1.00 17.55 | B_13 |
| MOTA | 2293 | CD | PRO | 87 | 46.565 | 56.165 | 4.146 | 1.00 10.17 | B_13 |
| ATOM | 2294 | CA | PRO | 87 | 46.228 | 58.041 | 5.628 | 1.00 32.09 | B_13 |
| MOTA | 2295 | CB | PRO | 87 | 44.961 | 57.720 | 4.819 | 1.00 18.55 | B_13 |
| ATOM | 2296 | CG | PRO | 87 | 45.115 | 56.277 | 4.481 | 1.00 18.86 | B_13 |
| ATOM | 2297 | С | PRO | 87 | 45.995 | 57.955 | 7.139 | 1.00 25.18 | B_13 |
| MOTA | 2298 | 0 | PRO | 87 | 46.284 | 56.919 | 7.752 | 1.00 18.18 | B_13 |
| MOTA | 2299 | N | PRO | 88 | 45.462 | 59.032 | 7.760 | 1.00 11.49 | B_13 |
| ATOM | 2300 | CD | PRO | 88 | 45.015 | 60.303 | 7.164 | 1.00 10.00 | B_13 |
| ATOM | 2301 | CA | PRO | 88 | 45.217 | 59.034 | 9.202 | 1.00 19.03 | B_13 |
| MOTA | 2302 | CB | PRO | 88 | 44.399 | 60.302 | 9.402 | 1.00 14.16 | B_13 |
| MOTA | 2303 | CG | PRO | 88 | 44.939 | 61.196 | 8.357 | 1.00 16.39 | B_13 |
| MOTA | 2304 | С | PRO | 88 | 44.500 | 57.787 | 9.733 | 1.00 25.43 | B_13 |
| MOTA | 2305 | 0 | PRO | 88 | 43.670 | 57.165 | 9.044 | 1.00 15.90 | B_13 |
| MOTA | 2306 | N | GLY | 89 | 44.865 | 57.422 | 10.955 | 1.00 26.28 | B_13 |
| MOTA | 2308 | CA | GLY | 89 | 44.299 | 56.264 | 11.606 | 1.00 25.32 | B_13 |
| MOTA | 2309 | С | GLY | 89 | 45.343 | 55.713 | 12.546 | 1.00 34.38 | B_13 |
| MOTA | 2310 | 0 | GLY | 89 | 46.485 | 56.164 | 12.498 | 1.00 23.28 | B_13 |
| MOTA | 2311 | N | PRO | 90 | 44.977 | 54.774 | 13.437 | 1.00 13.87 | B_13 |
| MOTA | 2312 | CD | PRO | 90 | 43.613 | 54.259 | 13.631 | 1.00 16.36 | B_13 |
| ATOM | 2313 | CA | PRO | 90 | 45.898 | 54.164 | 14.398 | 1.00 10.34 | B_13 |
| MOTA | 2314 | | PRO | 90 | 44.963 | 53.360 | 15.300 | 1.00 15.93 | B_13 |
| MOTA | . 2315 | CG | PRO | 90 | 43.870 | 52.975 | | 1.00 23.25 | B_13 |
| MOTA | 2316 | C | PRO | 90 | 46.942 | 53.299 | 13.711 | 1.00 18.38 | B_13 |
| MOTA | 2317 | 0 | PRO | 90 | 46.875 | 53.064 | 12.505 | 1.00 26.81 | B_13 |
| MOTA | 2318 | N | ASN | 91 | 47.903 | 52.831 | 14.502 | 1.00 26.63 | B_13 |
| ATOM ATOM | 2320 2321 | CA CB | asn Asn | 91 91 | 49.022 48.740 | 52.010 50.500 | 14.033 | 1.00 21.91 | B_13 |
| MOTA | 2322 | CG | ASN | 91 | 47.437 | 50.300 | 14.081 13.448 | 1.00 18.89 | B_13 |
| MOTA | 2323 | | ASN | 91 | 47.335 | 50.117 | 12.237 | 1.00 22.49 1.00 29.37 | B_13 |
| MOTA | 2324 | ND2 | | 91 | 46.438 | 49.858 | 14.273 | 1.00 28.01 | B_13 B_13 |
| ATOM | 2327 | C | ASN | 91 | 49.656 | 52.438 | 12.721 | 1.00 20.07 | B_13 |
| ATOM | 2328 | Ö | ASN | 91 | 50.301 | 53.479 | 12.681 | 1.00 21.24 | B_13 |
| MOTA | 2329 | N | TYR | 92 | 49.423 | 51.716 | 11.633 | 1.00 20.15 | B_13 |
| MOTA | 2331 | CA | TYR | 92 | 50.052 | 52.081 | 10.367 | 1.00 18.70 | B_13 |
| MOTA | 2332 | CB | TYR | 92 | 49.905 | 50.953 | 9.344 | 1.00 14.48 | B_13 |
| ATOM | 2333 | CG | TYR | 92 | 50.906 | 49.821 | 9.567 | 1.00 24.41 | B_13 |
| MOTA | 2334 | CD1 | TYR | 92 | 52.266 | 50.003 | 9.287 | 1.00 27.39 | B_13 |
| MOTA | 2335 | CE1 | | 92 | 53.198 | 48.979 | 9.471 | 1.00 18.14 | B_13 |
| MOTA | 2336 | CD2 | | 92 | 50.499 | 48.571 | 10.044 | 1.00 28.07 | B_13 |
| ATOM | 2337 | CE2 | | 92 | 51.427 | 47.529 | 10.230 | 1.00 36.50 | B_13 |
| MOTA | 2338 | CZ | TYR | 92 | 52.778 | 47.741 | 9.940 | 1.00 43.64 | B_13 |
| ATOM | 2339 | OH | TYR | 92 | 53.694 | 46.710 | 10.105 | 1.00 32.21 | B_13 |
| MOTA | 2341 | C | TYR | 92 | 49.633 | 53.431 | 9.797 | 1.00 21.78 | B_13 |
| MOTA | 2342 | 0 | TYR | 92 | 50.384 | 54.049 | 9.040 | 1.00 12.55 | B_13 |
| MOTA | 2343 | N | GLY | 93 | 48.464 | 53.916 | 10.198 | 1.00 15.83 | B_13 |
| ATOM | 2345 2346 | CA | GLY | 93 | 48.015 | 55.216 | 9.732 | | B_13 |
| MOTA MOTA | 2347 | C O | GLY | 93 | 48.971 | | 10.134 | 1.00 18.60 | B_13 |
| ATOM | 2348 | N | GLY | 93 | 49.561 | 56.300 | 11.227 | 1.00 22.00 | B_13 |
| ATOM | 2350 | CA | GLY GLY | 94 94 | 49.205 | 57.258 | 9.216 | 1.00 10.27 | B_13 |
| ATOM | 2351 | C | GLY | 94 | 50.099 | 58.365 | 9.492 | | B_13 |
| ATOM | 2352 | 0 | GLY | 94 94 | 51.567 52.334 | 58.061 58.967 | 9.234 | | B_13 |
| ATOM | 2353 | N | ASP | 9 4 95 | 52.334 | | 8.938 9.351 | | B_13 |
| ATOM | 2355 | CA | ASP | | 53.386 | | 9.331 | | B_13 B_13 |
| ATOM | 2356 | CB | ASP | 95 95 | 53.560 | 54.986 | 9.134 | | B_13 B_13 |
| ATOM | 2357 | CG | ASP | 95 | 53.346 | | 10.900 | | B_13 |
| ATOM | 2358 | ODI | | | 53.627 | | 11.297 | | B_13 |
| ATOM | 2359 | OD2 | | | 52.835 | 55.488 | 11.656 | 1.00 14.66 | B_13 |
| ATOM | 2360 | C | ASP | | 53.896 | | 7.733 | | B_13 |
| ATOM | 2361 | Ö | ASP | | 53.162 | | 6.746 | | B_13 |
| ATOM | 2362 | N | ALA | | 55.166 | | 7.662 | | B_13 |
| MOTA | 2364 | CA | ALA | | 55.803 | | 6.400 | | B_13 |

| ATOM | 2365 | СВ | ALA | 96 | 56.098 | 59.095 | 6.379 | 1.00 22.61 | n 13 |
|------|------|-----|------|-------|--------|---------|--------|------------|------|
| | | | | | 57.088 | 56.784 | | | B_13 |
| ATOM | 2366 | | ALA | 96 | | • | 6.204 | 1.00 25.63 | B_13 |
| ATOM | 2367 | | ALA | 96 | 57.948 | 56.724 | | 1.00 12.54 | B_13 |
| ATOM | 2368 | | HIS | 97 | 57.211 | 56.166 | 5.035 | 1.00 13.27 | B_13 |
| MOTA | 2370 | CA | HIS | 97 | 58.375 | 55.357 | 4.730 | 1.00 25.28 | B_13 |
| MOTA | 2371 | CB | HIS | 97 | 57.955 | 53.905 | 4.464 | 1.00 10.00 | B_13 |
| MOTA | 2372 | CG | HIS | 97 | 57.264 | 53.257 | 5.624 | 1.00 12.02 | B_13 |
| ATOM | 2373 | _ | HIS | 97 | 57.214 | 53.603 | 6.929 | 1.00 10.00 | B_13 |
| ATOM | 2374 | | HIS | 97 | 56.516 | 52.104 | 5.499 | 1.00 12.91 | |
| | | | | | | | | | B_13 |
| MOTA | 2375 | | HIS | 97 | 56.038 | 51.770 | 6.688 | 1.00 10.00 | B_13 |
| MOTA | 2376 | | HIS | 97 | 56.445 | 52.664 | 7.571 | 1.00 10.64 | B_13 |
| MOTA | 2378 | C | HIS | 97 | 59.069 | 55.959 | 3.520 | 1.00 13.82 | B_13 |
| ATOM | 2379 | 0 | HIS | 97 | 58.415 | 56.273 | 2.517 | 1.00 12.27 | B_13 |
| MOTA | 2380 | N | PHE | 98 | 60.379 | 56.154 | 3.647 | 1.00 10.67 | B_13 |
| MOTA | 2382 | | PHE | 98 | 61.224 | 56.718 | 2.595 | 1.00 15.67 | B_13 |
| ATOM | 2383 | | PHE | 98 | 61.970 | 57.938 | 3.156 | 1.00 10.76 | B_13 |
| ATOM | 2384 | | PHE | 98 | 61.055 | 59.025 | 3.627 | | |
| | | | | | | | | 1.00 17.93 | B_13 |
| ATOM | 2385 | | PHE | 98 | 60.730 | 60:082 | 2.786 | 1.00 18.92 | B_13 |
| ATOM | 2386 | | PHE | 98 | 60.476 | 58.974 | 4.893 | 1.00 14.14 | B_13 |
| ATOM | 2387 | | PHE | 98 | 59.833 | 61.066 | 3.201 | 1.00 22.42 | B_13 |
| ATOM | 2388 | CE2 | PHE | 98 | 59.574 | 59.962 | 5.315 | 1.00 10.00 | B_13 |
| MOTA | 2389 | CZ | PHE. | 98 | 59.257 | 61.002 | 4.469 | 1.00 10.00 | B_13 |
| MOTA | 2390 | С | PHE | 98 | 62.218 | 55.669 | 2.064 | 1.00 26.64 | B_13 |
| MOTA | 2391 | 0 | PHE | 98 | 62.882 | 54.969 | 2.851 | 1.00 13.27 | B_13 |
| MOTA | 2392 | N | ASP | 99 | 62.331 | 55.577 | 0.738 | 1.00 12.24 | B_13 |
| ATOM | 2394 | CA | ASP | 99 | 63.229 | | | | |
| | | | | | | 54.612 | 0.102 | 1.00 10.00 | B_13 |
| ATOM | 2395 | | ASP | 99 | 62.884 | 54.471 | -1.385 | 1.00 10.00 | B_13 |
| ATOM | 2396 | CG | ASP | 99 | 63.615 | 53.311 | -2.067 | 1.00 22.86 | B_13 |
| MOTA | 2397 | OD1 | | 99 | 63.170 | 52.890 | -3.160 | 1.00 11.60 | B_13 |
| MOTA | 2398 | OD2 | ASP | 99 | 64.624 | 52.806 | -1.528 | 1.00 21.20 | B_13 |
| MOTA | 2399 | C | ASP | 99 | 64.677 | 55.046 | 0.264 | 1.00 12.66 | B_13 |
| ATOM | 2400 | 0 | ASP | 99 | 65.121 | 56.010 | -0.366 | 1.00 18.37 | B_13 |
| MOTA | 2401 | N | ASP | 100 | 65.439 | 54.289 | 1.046 | 1.00 12.86 | B_13 |
| ATOM | 2403 | CA | ASP | 100 | 66.833 | 54.642 | 1.260 | 1.00 14.46 | B_13 |
| ATOM | 2404 | CB | ASP | 100 | | | | | |
| | | | | | 67.308 | 54.271 | 2.660 | 1.00 17.70 | B_13 |
| MOTA | 2405 | CG | ASP | 100 | 68.006 | 55.437 | 3.358 | 1.00 16.15 | B_13 |
| MOTA | 2406 | | ASP | 100 | 68.091 | 55.447 | 4.602 | 1.00 15.74 | B_13 |
| MOTA | 2407 | | ASP | 100 | 68.470 | 56.354 | 2.655 | 1.00 27.08 | B_13 |
| MOTA | 2408 | С | ASP | 100 | 67.793 | 54.171 | 0.179 | 1.00 13.66 | B_13 |
| ATOM | 2409 | 0 | ASP | 100 | 68.961 | 53.932 | 0.416 | 1.00 19.54 | B_13 |
| ATOM | 2410 | N | ASP | 101 | 67.254 | 53.954 | -1.010 | 1.00 12.83 | B_13 |
| ATOM | 2412 | CA | ASP | 101 | 68.074 | 53.590 | -2.164 | 1.00 10.00 | B_13 |
| MOTA | 2413 | CB | ASP | 101 | 67.471 | 52.413 | -2.933 | 1.00 10.00 | |
| ATOM | 2414 | | ASP | 101 | | | | | B_13 |
| | | | | | 67.997 | 51.065 | -2.449 | 1.00 16.87 | B_13 |
| MOTA | 2415 | | ASP | 101 | 67.232 | 50.089 | -2.458 | 1.00 19.89 | B_13 |
| ATOM | 2416 | | ASP | 101 | 69.184 | 50.968 | -2.066 | 1.00 18.51 | B_13 |
| MOTA | 2417 | C | ASP | 101 | 68.108 | 54.858 | -3.029 | 1.00 26.72 | B_13 |
| ATOM | 2418 | 0 | ASP | 101 | 68.602 | 54.853 | -4.172 | 1.00 12.11 | B_13 |
| ATOM | 2419 | N | GLU | 102 | 67.500 | 55.922 | -2.496 | 1.00 13.76 | B_13 |
| MOTA | 2421 | CA | GLŲ | 102 | 67.462 | 57.217 | -3.161 | 1.00 12.54 | B_13 |
| ATOM | 2422 | CB | GLU | 102 | 66.135 | 57.958 | -2.916 | 1.00 13.01 | B_13 |
| ATOM | 2423 | CG | GLU | 102 | 64.873 | 57.257 | -3.381 | 1.00 15.50 | B_13 |
| ATOM | 2424 | CD | GLU | 102 | 64.973 | 56.707 | -4.791 | 1.00 29.02 | B_13 |
| MOTA | 2425 | | GLU | 102 | 65.640 | 57.307 | -5.665 | | |
| ATOM | 2426 | OE2 | | 102 | | | | 1.00 12.78 | B_13 |
| ATOM | 2427 | C | GLU | 102 | 64.399 | 55.635 | -5.021 | 1.00 12.36 | B_13 |
| | | | | | 68.544 | | -2.505 | | B_13 |
| MOTA | 2428 | 0 | GLU | 102 | 68.939 | 57.760 | -1.371 | 1.00 10.00 | B_13 |
| ATOM | 2429 | N | THR | 103 | 69.030 | 59.039 | -3.228 | 1.00 19.38 | B_13 |
| ATOM | 2431 | CA | THR | 103 | 70.021 | 59.957 | -2.693 | 1.00 16.49 | B_13 |
| ATOM | 2432 | CB | THR | 103 | 70.973 | 60.490 | -3.801 | 1.00 19.31 | B_13 |
| MOTA | 2433 | OG1 | THR | 103 | 71.661 | 59.384 | -4.399 | 1.00 25.44 | B_13 |
| ATOM | 2435 | CG2 | THR | 103 | 72.006 | 61.462 | -3.212 | | B_13 |
| ATOM | 2436 | C | THR | 103 | 69.180 | 61.104 | -2.141 | 1.00 12.91 | B_13 |
| MOTA | 2437 | ō | THR | 103 | 68.414 | 61.727 | -2.867 | | B_13 |
| ATOM | 2438 | N | TRP | 104 | 69.252 | | -0.842 | | |
| ATOM | 2440 | CA | TRP | 104 | | | | | B_13 |
| | | | | | 68.497 | | -0.237 | | B_13 |
| MOTA | 2441 | CB | TRP | 104 | 67.852 | 61.902 | 1.063 | 1.00 22.66 | B_13 |
| ATOM | 2442 | CG | TRP | 104 | 66.837 | | 0.870 | | B_13 |
| MOTA | 2443 | | TRP | 104 | 65.505 | | 0.347 | 1.00 27.35 | B_13 |
| ATOM | 2444 | | TRP | 104 | 64.936 | | 0.287 | 1.00 12.61 | B_13 |
| MOTA | 2445 | CE3 | TRP | 104 | 64.741 | | -0.079 | | B_13 |
| MOTA | 2446 | CD1 | TRP | 104 | 67.013 | | 1.108 | _ | B_13 |
| ATOM | 2447 | | TRP | 104 | 65.876 | | 0.755 | | B_13 |
| ATOM | 2449 | | TRP | 104 | 63.632 | | -0.186 | | B_13 |
| MOTA | 2450 | | TRP | 104 | 63.445 | | -0.549 | | |
| ATOM | 2451 | CH2 | | 104 | 62.904 | | | • | B_13 |
| ATOM | 2452 | C | TRP | 104 | 69.416 | | -0.598 | | B_13 |
| | | _ | -115 | + U = | 03.470 | 010.010 | 0.033 | 1.00 16.43 | B_13 |

| | 2153 | _ | | | | | | | |
|------|------|-----|-----|-----|--------|--------|--------|------------|------|
| MOTA | 2453 | 0 | TRP | 104 | 70.520 | 63.380 | 0.526 | 1.00 11.13 | B_13 |
| ATOM | 2454 | N | THR | 105 | 68.960 | 64.775 | -0.322 | 1.00 19.48 | B_13 |
| ATOM | 2456 | CA | THR | 105 | 69.716 | 66.015 | -0.097 | 1.00 10.40 | B_13 |
| MOTA | 2457 | CB | THR | 105 | 70.153 | 66.749 | -1.398 | 1.00 10.00 | B_13 |
| MOTA | 2458 | OG1 | THR | 105 | 69.305 | 66.401 | -2.501 | 1.00 18.53 | B_13 |
| MOTA | 2460 | CG2 | THR | 105 | 71.596 | 66.484 | -1.709 | 1.00 34.62 | B_13 |
| ATOM | 2461 | С | THR | 105 | 68.904 | 67.062 | 0.641 | 1.00 20.82 | B_13 |
| MOTA | 2462 | 0 | THR | 105 | 67.686 | 66.952 | 0.768 | 1.00 15.93 | B_13 |
| MOTA | 2463 | N | SER | 106 | 69.621 | 68.073 | 1.125 | 1.00 38.37 | B_13 |
| MOTA | 2465 | CA | SER | 106 | 69.029 | 69.222 | 1.791 | 1.00 20.77 | B_13 |
| MOTA | 2466 | CB | SER | 106 | 69.979 | 69.778 | | | |
| | 2467 | OG | | 106 | | | 2.862 | 1.00 17.95 | B_13 |
| MOTA | | | SER | | 70.281 | 68.825 | 3.864 | 1.00 29.88 | B_13 |
| ATOM | 2469 | C | SER | 106 | 68.889 | 70.245 | 0.657 | 1.00 19.23 | B_13 |
| ATOM | 2470 | 0 | SER | 106 | 68.202 | 71.260 | 0.782 | 1.00 21.34 | B_13 |
| MOTA | 2471 | N | SER | 107 | 69.577 | 69.981 | -0.450 | 1.00 18.73 | B_13 |
| ATOM | 2473 | CA | SER | 107 | 69.533 | 70.884 | -1.592 | 1.00 20.92 | B_13 |
| MOTA | 2474 | CB | SER | 107 | 70.945 | 71.380 | -1.927 | 1.00 19.84 | B_13 |
| ATOM | 2475 | OG | SER | 107 | 71.556 | 71.957 | -0.788 | 1.00 27.31 | B_13 |
| MOTA | 2477 | С | SER | 107 | 68.848 | 70.284 | -2.828 | 1.00 18.68 | B_13 |
| MOTA | 2478 | 0 | SER | 107 | 67.660 | 69.953 | -2.771 | 1.00 21.51 | B_13 |
| MOTA | 2479 | N | SER | 108 | 69.623 | 70.038 | -3.888 | 1.00 18.53 | B_13 |
| ATOM | 2481 | CA | SER | 108 | 69.091 | 69.544 | -5.152 | 1.00 16.21 | B_13 |
| ATOM | 2482 | CB | SER | 108 | 69.285 | 70.632 | -6.205 | 1.00 29.10 | B_13 |
| ATOM | 2483 | OG | SER | 108 | 70.665 | 70.969 | -6.271 | 1.00 21.47 | B_13 |
| ATOM | 2485 | C | SER | 108 | 69.645 | 68.260 | -5.745 | 1.00 17.68 | B_13 |
| ATOM | 2486 | ŏ | SER | 108 | 68.964 | 67.618 | -6.541 | 1.00 19.67 | B_13 |
| ATOM | 2487 | Ň | LYS | 109 | 70.895 | 67.919 | -5.448 | 1.00 11.70 | |
| MOTA | 2489 | CA | LYS | 109 | 71.468 | | | | B_13 |
| MOTA | 2490 | CB | LYS | | | 66.721 | -6.047 | 1.00 10.00 | B_13 |
| | | | | 109 | 72.994 | 66.748 | -5.989 | 1.00 18.86 | B_13 |
| MOTA | 2491 | CG | LYS | 109 | 73.657 | 65.833 | -7.013 | 1.00 16.33 | B_13 |
| MOTA | 2492 | CD | LYS | 109 | 75.143 | 65.726 | -6.740 | 1.00 11.58 | B_13 |
| MOTA | 2493 | CE | LYS | 109 | 75.787 | 64.655 | -7.606 | 1.00 27.43 | B_13 |
| MOTA | 2494 | NZ | LYS | 109 | 77.218 | 64.492 | -7.251 | 1.00 35.03 | B_13 |
| MOTA | 2498 | С | LYS | 109 | 70.916 | 65.428 | -5.444 | 1.00 29.39 | B_13 |
| MOTA | 2499 | .0 | LYS | 109 | 71.432 | 64.905 | -4.449 | 1.00 29.95 | B_13 |
| MOTA | 2500 | N | GLY | 110 | 69.852 | 64.922 | -6.055 | 1.00 14.77 | B_13 |
| MOTA | 2502 | CA | GLY | 110 | 69.227 | 63.705 | -5.576 | 1.00 24.08 | B_13 |
| MOTA | 2503 | С | GLY | 110 | 67.793 | 64.105 | -5.342 | 1.00 20.25 | B_13 |
| ATOM | 2504 | O | GLY | 110 | 67.203 | 64.737 | -6.198 | 1.00 16.21 | B_13 |
| ATOM | 2505 | N | TYR | 111 | 67.248 | 63.772 | -4.182 | 1.00 10.00 | B_13 |
| MOTA | 2507 | CA | TYR | 111 | 65.879 | 64.130 | -3.845 | 1.00 10.00 | |
| ATOM | 2508 | CB | TYR | 111 | | | | | B_13 |
| | 2509 | | | | 65.030 | 62.868 | -3.688 | 1.00 22.46 | B_13 |
| MOTA | | CG | TYR | 111 | 64.676 | 62.244 | | 1.00 10.83 | B_13 |
| MOTA | 2510 | CD1 | | 111 | 65,380 | 61.155 | -5.483 | 1.00 25.38 | B_13 |
| ATOM | 2511 | CE1 | | 111 | 65.068 | 60.592 | -6.720 | 1.00 18.68 | B_13 |
| ATOM | 2512 | CD2 | | 111 | 63.646 | 62.769 | | 1.00 16.02 | B_13 |
| MOTA | 2513 | CE2 | | 111 | 63.328 | | -7.013 | 1.00 31.72 | B_13 |
| ATOM | 2514 | CZ | TYR | 111 | 64.041 | 61.131 | -7.473 | 1.00 23.68 | B_13 |
| ATOM | 2515 | OH | TYR | 111 | 63.711 | 60.550 | -8.666 | 1.00 20.96 | B_13 |
| ATOM | 2517 | C | TYR | 111 | 65.856 | 64.944 | -2.553 | 1.00 22.83 | B_13 |
| MOTA | 2518 | 0 | TYR | 111 | 66.410 | 64.518 | -1.538 | 1.00 11.66 | B_13 |
| ATOM | 2519 | N | ASN | 112 | 65.278 | 66.140 | -2.611 | 1.00 17.47 | B_13 |
| MOTA | 2521 | CA | ASN | 112 | 65.180 | 67.006 | -1.431 | 1.00 15.77 | B_13 |
| MOTA | 2522 | CB | ASN | 112 | 64.658 | 68.401 | -1.817 | 1.00 15.93 | B_13 |
| MOTA | 2523 | CG | ASN | 112 | 64.694 | 69.384 | -0.657 | 1.00 10.00 | B_13 |
| MOTA | 2524 | OD1 | ASN | 112 | 63.757 | 69.465 | 0.132 | 1.00 15.33 | B_13 |
| ATOM | 2525 | ND2 | ASN | | 65.754 | 70.180 | -0.586 | 1.00 13.70 | B_13 |
| MOTA | 2528 | C | ASN | 112 | 64.214 | 66.329 | -0.472 | 1.00 17.73 | B_13 |
| MOTA | 2529 | 0 | ASN | 112 | 63.007 | 66.243 | -0.737 | 1.00 12.61 | B_13 |
| ATOM | 2530 | N | LEU | | 64.755 | 65.830 | 0.630 | 1.00 16.28 | B_13 |
| ATOM | 2532 | CA | LEU | 113 | 63.962 | 65.121 | | | |
| ATOM | 2533 | CB | LEU | 113 | | | 1.619 | 1.00 15.93 | B_13 |
| | | | | | 64.841 | 64.703 | 2.804 | 1.00 11.93 | B_13 |
| ATOM | 2534 | CG | LEU | | 64.719 | 63.352 | | 1.00 17.15 | B_13 |
| ATOM | 2535 | | LEU | | 65.002 | 63.640 | 4.987 | 1.00 10.00 | B_13 |
| ATOM | 2536 | CD2 | | | 63.370 | 62.667 | 3.362 | 1.00 16.08 | B_13 |
| MOTA | 2537 | С | LEU | | 62.802 | 65.994 | 2.085 | 1.00 14.61 | B_13 |
| MOTA | 2538 | 0 | LEU | | 61.673 | 65:528 | 2.161 | 1.00 17.98 | B_13 |
| ATOM | 2539 | N | PHE | | 63.073 | 67.267 | 2.346 | 1.00 16.81 | B_13 |
| ATOM | 2541 | CA | PHE | | 62.056 | 68.212 | 2.791 | 1.00 15.65 | B_13 |
| MOTA | 2542 | CB | PHE | | 62.638 | 69.630 | 2.888 | 1.00 22.16 | B_13 |
| MOTA | 2543 | CG | PHE | | 61.596 | 70.714 | 2.882 | 1.00 12.27 | B_13 |
| ATOM | 2544 | CD1 | | | 60.804 | 70.952 | 4.004 | 1.00 19.93 | B_13 |
| ATOM | 2545 | | PHE | | 61.378 | | 1.746 | 1.00 13.56 | B_13 |
| ATOM | 2546 | | PHE | | 59.813 | 71.932 | 3.984 | 1.00 13.38 | D_13 |
| ATOM | 2547 | CE2 | | | 60.398 | 72.441 | | | B_13 |
| ATOM | 2548 | CZ | PHE | | | | 1.726 | 1.00 13.79 | B_13 |
| | | | | | 59.615 | | 2.848 | 1.00 10.70 | B_13 |
| MOTA | 2549 | С | PHE | 114 | 60.860 | 68.220 | 1.842 | 1.00 19.55 | B_13 |

| ATOM | 2550 | 0 | PHE | 114 | 59.714 | 68.156 | 2.285 | 1.00 15.97 | B_13 |
|--------------|--------------|-----------|------------|------------|------------------|------------------|------------------|--------------------------|--------------|
| ATOM | 2551 | N | LEU | 115 | 61.135 | 68.309 | 0.543 | 1.00 13.35 | B_13 |
| ATOM | 2553 | | LEU | 115 | 60.096 | 68.323 | -0.485 | 1.00 17.91 | B_13 |
| ATOM | 2554 | CB | LEU | 115 | 60.741 | 68.462 | -1.868 | 1.00 24.65 | B_13 |
| ATOM | 2555 | CG | LEU | 115 | 60.501 | 69.739 | -2.679 | 1.00 22.70 | B_13 |
| ATOM | 2556 | CD1 | LEU | 115 | 61.033 | 70.939 | -1.943 | 1.00 17.98 | B_13 |
| ATOM | 2557 | | LEU | 115 | 61.148 | 69.624 | -4.048 | 1.00 28.50 | B_13 |
| ATOM | 2558 | C | LEU | 115 | 59.235 | 67.042 | -0.443 | 1.00 21.61 | B_13 |
| MOTA | 2559 | 0 | LEU | 115 | 58.002 | 67.093 | -0.344 | 1.00 13.99 | B_13 |
| MOTA | 2560 | N | VAL | 116 | 59.898 | 65.895 | -0.511 | 1.00 11.14 | B_13 |
| MOTA | 2562 | CA | VAL | 116 | 59.199 | 64.616 | -0.482 | 1.00 22.27 | B_13 |
| MOTA | 2563 2564 | CB CG1 | VAL VAL | 116 116 | 60.163 | 63.421 62.086 | -0.772 -0.629 | 1.00 17.40 | B_13 |
| MOTA MOTA | 2565 | CG2 | VAL | 116 | 59.437 60.741 | 63.534 | -0.029 | 1.00 23.09 1.00 12.16 | B_13 B_13 |
| ATOM | 2566 | C | VAL | 116 | 58.502 | 64.414 | 0.864 | 1.00 10.00 | B_13 |
| ATOM | 2567 | Ö | VAL | 116 | 57.368 | 63.950 | 0.911 | 1.00 16.18 | B_13 |
| ATOM | 2568 | N | ALA | 117 | 59.153 | 64.803 | 1.954 | 1.00 10.00 | B_13 |
| MOTA | 2570 | CA | ALA | 117 | 58.585 | 64.640 | 3.297 | 1.00 19.50 | B_13 |
| MOTA | 2571 | CB | ALA | 117 | 59.608 | 64.995 | 4.352 | 1.00 11.81 | B_13 |
| MOTA | 2572 | C | ALA | 117 | 57.309 | 65.455 | 3.505 | 1.00 30.87 | B_13 |
| ATOM | 2573 | 0 | ALA | 117 | 56.327 | 64.955 | 4.053 | 1.00 10.00 | B_13 |
| ATOM | 2574 | N | ALA | 118 | 57.322 | 66.714 | 3.087 | 1.00 24.62 | B_13 |
| MOTA | 2576 | CA | ALA | 118 | 56.140 | 67.553 | 3.222 | 1.00 20.76 | B_13 |
| MOTA MOTA | 2577 2578 | CB C | ALA ALA | 118 118 | 56.407 54.968 | 68.917 66.894 | 2.654 2.485 | 1.00 16.19 1.00 20.54 | B_13 |
| ATOM | 2579 | Ö | ALA | 118 | 53.843 | 66.889 | 2.485 | 1.00 20.34 | B_13 B_13 |
| ATOM | 2580 | N | HIS | 119 | 55.255 | 66.315 | 1.321 | 1.00 10.00 | B_13 |
| MOTA | 2582 | CA | HIS | 119 | 54.259 | 65.647 | 0.489 | 1.00 17.27 | B_13 |
| MOTA | 2583 | CB | HIS | 119 | 54.909 | 65.263 | -0.860 | 1.00 11.16 | B_13 |
| MOTA | 2584 | | HIS | 119 | 54.006 | 64.530 | -1.813 | 1.00 26.59 | ' B_13 |
| MOTA | 2585 | CD2 | HIS | 119 | 53.377 | 63.335 | -1.706 | 1.00 16.63 | B_13 |
| ATOM | 2586 | | HIS | 119 | 53.723 | 64.995 | -3.085 | 1.00 12.44 | B_13 |
| MOTA | 2588 | | HIS | 119 | 52.961 | 64.124 | | 1.00 14.58 | B_13 |
| ATOM | 2589 | | HIS | 119 | 52.734 | 63.101 | -2.901 | 1.00 26.44 | B_13 |
| ATOM | 2590 | C | HIS | 119 | 53.722 | 64.419 | 1.227 | 1.00 17.00 | B_13 |
| ATOM | 2591 | 0 | HIS | 119 | 52.510 | 64.218 | | 1.00 17.01 | _ |
| MOTA MOTA | 2592 2594 | N CA | GLU GLU | 120 120 | 54.626 54.231 | 63.607 | 1.751 | 1.00 10.31 | B_13 |
| MOTA | 2595 | CB | GLU | 120 | 54.231 55.463 | 62.401 61.627 | 2.466 2.961 | 1.00 12.32 1.00 15.34 | B_13 |
| MOTA | 2596 | CG | GLU | 120 | 56.354 | 61.078 | 1.848 | 1.00 10.00 | B_13 B_13 |
| ATOM | 2597 | CD | GLU | 120 | 55.574 | 60.260 | 0.867 | 1.00 18.64 | B_13 |
| ATOM | 2598 | OE1 | | 120 | 55.598 | 60.565 | -0.348 | 1.00 18.08 | B_13 |
| ATOM | 2599 | OE2 | GLU | 120 | 54.920 | 59.308 | | .1.00 14.49 | |
| ATOM | 2600 | C | GĽU | 120 | 53.347 | 62.777 | 3.635 | 1.00 12.41 | B_13 |
| MOTA | 2601 | 0 | GLU | 120 | 52.323 | 62.130 | 3.888 | 1.00 26.62 | B_13 |
| MOTA | 2602 | N | PHE | 121 | 53.750 | 63.813 | | | B_13 |
| MOTA | 2604 | CA | PHE | 121 | 52.993 | 64.286 | 5.506 | | - |
| MOTA MOTA | 2605 2606 | CB CG | PHE | 121 121 | 53.780 55.057 | 65.344 64.827 | 6.270 6.852 | 1.00 20.10 1.00 24.55 | B_13 |
| ATOM | 2607 | CD1 | | 121 | 56.037 | 65.700 | 7.292 | | B_13 B_13 |
| ATOM | 2608 | | PHE | 121 | 55.292 | 63.454 | | 1.00 23.62 | B_13 |
| ATOM | 2609 | CE1 | | 121 | 57.247 | 65.212 | 7.813 | 1.00 18.59 | |
| MOTA | 2610 | CE2 | | 121 | 56.488 | 62.954 | 7.448 | | B_13 |
| MOTA | 2611 | CZ | PHE | 121 | 57.472 | 63.834 | 7.888 | 1.00 25.40 | B_13 |
| ATOM | 2612 | C | PHE | 121 | 51.607 | 64.791 | 5.110 | | B_13 |
| MOTA | 2613 | 0 | PHE | 121 | 50.676 | 64.760 | 5.921 | | B_13 |
| MOTA | 2614 | N | GLY | | 51.471 | 65.238 | 3.864 | | B_13 |
| MOTA | 2616 2617 | CA | GLY | 122 122 | 50.175 | 65.664 | 3.380 | | B_13 |
| ATOM ATOM | 2618 | C O | GLY GLY | 122 | 49.284 48.113 | 64.427 64.483 | 3.381 | 1.00 13.71 | B_13 |
| ATOM | 2619 | N | HIS | 123 | 49.859 | 63.284 | 3.753 3.016 | 1.00 13.74 1.00 16.90 | B_13 B_13 |
| ATOM | 2621 | CA | HIS | 123 | 49.126 | 62.009 | 3.008 | | B_13 |
| ATOM | 2622 | CB | HIS | 123 | 49.918 | 60.918 | 2.279 | | B_13 |
| ATOM | 2623 | CG | HIS | 123 | 49.945 | 61.084 | 0.794 | | B_13 |
| MOTA | 2624 | CD2 | | 123 | 50.889 | 60.764 | -0.119 | | B_13 |
| MOTA | 2625 | ND | HIS | 123 | 48.887 | 61.618 | | | B_13 |
| MOTA | 2627 | | HIS | 123 | 49.176 | 61.621 | -1.195 | 1.00 16.02 | B_13 |
| ATOM | 2628 | NE2 | | 123 | 50.386 | 61.108 | -1.353 | | B_13 |
| ATOM | 2629 | C | HIS | 123 | 48.864 | 61.562 | | | B_13 |
| MOTA | 2630 | | HIS | | 47.744 | 61.179 | | | B_13 |
| MOTA MOTA | 2631 2633 | N | SER | | 49.904 | 61.627 | | | B_13 |
| ATOM | 2634 | CA CB | SER SER | | 49.813 51.131 | 61.270 61.582 | - | | B_13 |
| ATOM | 2635 | | SER | | 52.221 | 60.837 | | | B_13 B_13 |
| MOTA | 2637 | C | SER | | 48.703 | 62.102 | 7.335 | | B_13 |
| MOTA | 2638 | | SER | | 48.061 | 61.677 | | | B_13 |
| MOTA | 2639 | N | LEU | 125 | 48.481 | 63.300 | | | B_13 |
| | | | | | | | | | |

| ATOM | 2641 | CA | LEU | 125 | 47.439 | 64.133 | 7.387 | 1.00 24.62 | ' D 12 |
|------|------|-----|-----|-----|--------|--------|----------------------|------------|-------------|
| | | | | | | | | | B_13 |
| MOTA | 2642 | CB | LEU | 125 | 47.893 | 65.592 | 7.436 | 1.00 20.76 | B_13 |
| ATOM | 2643 | CG | LEU | 125 | 49.076 | 65.849 | 8.383 | 1.00 14.66 | B_13 |
| ATOM | 2644 | CD1 | | 125 | 49.739 | 67.159 | 8.064 | 1.00 16.16 | |
| | | | | | | | | | B_13 |
| ATOM | 2645 | CD2 | LEU | 125 | 48.610 | 65.811 | 9.822 | 1.00 16.44 | B_13 |
| ATOM | 2646 | С | LEU | 125 | 46.058 | 63.966 | 6.724 | 1.00 24.77 | B_13 |
| ATOM | 2647 | | | | | | | | |
| | | | LEU | 125 | 45.066 | 64.528 | 7.195 | 1.00 15.63 | B_13 |
| MOTA | 2648 | N | GLY | 126 | 45.988 | 63.192 | 5.644 | 1.00 17.38 | B_13 |
| ATOM | 2650 | | GLY | 126 | 44.700 | | • | | _ |
| | | | | | | 62.968 | 5.001 | 1.00 22.41 | B_13 |
| ATOM | 2651 | C | GLY | 126 | 44.453 | 63.487 | 3.603 | 1.00 13.20 | B_13 |
| MOTA | 2652 | | GLY | 126 | 43.349 | 63.366 | 3.096 | 1.00 20.86 | |
| | | | | | | | | | B_13 |
| MOTA | 2653 | N | LEU | 127 | 45.452 | 64.079 | 2.972 | 1.00 12.39 | B_13 |
| MOTA | 2655 | CA | LEU | 127 | 45.267 | 64.592 | 1.617 | 1.00 11.56 | B_13 |
| | | | | | | | | | |
| ATOM | 2656 | CB | LEU | 127 | 45.965 | 65.947 | 1.467 | 1.00 19.19 | B_13 |
| MOTA | 2657 | CG | LEU | 127 | 45.300 | 67.206 | 2.039 | 1.00 14.42 | B_13 |
| | 2658 | | | | | | | | |
| MOTA | | | LEU | 127 | 44.875 | 67.030 | 3.496 | 1.00 32.31 | B_13 |
| MOTA | 2659 | CD2 | LEU | 127 | 46.288 | 68.374 | 1.912 | 1.00 25.45 | B_13 |
| ATOM | 2660 | C | LEU | 127 | 45.770 | 63.619 | 0.550 | 1.00 26.54 | |
| | | | | | | | | | B_13 |
| MOTA | 2661 | 0 | LEU | 127 | 46.920 | 63.156 | 0.601 | 1.00 18.76 | B_13 |
| MOTA | 2662 | N | ASP | 128 | 44.908 | 63.285 | -0.407 | 1.00 28.54 | B_13 |
| | | | | | | | | | |
| MOTA | 2664 | CA | ASP | 128 | 45.292 | 62.376 | -1.480 | 1.00 10.89 | B_13 |
| MOTA | 2665 | CB | ASP | 128 | 44.059 | 61.762 | -2.136 | 1.00 15.95 | B_13 |
| MOTA | 2666 | CG | ASP | 128 | 44.351 | 60.430 | -2.794 | | 2723 |
| | | | | | | | | 1.00 23.44 | B_13 |
| ATOM | 2667 | OD1 | ASP | 128 | 43.377 | 59.735 | -3.164 | 1.00 41.43 | B_13 |
| ATOM | 2668 | OD2 | ASP | 128 | 45.541 | 60.059 | -2.918 | 1.00 18.12 | |
| | | | | | | | | | B_13 |
| ATOM | 2669 | С | ASP | 128 | 46.060 | 63.203 | -2.502 | 1.00 25.34 | B_13 |
| ATOM | 2670 | 0 | ASP | 128 | 46.489 | 64.308 | -2.213 | 1.00 16.36 | B_13 |
| | | _ | | | | | | | |
| MOTA | 2671 | N | HIS | 129 | 46.283 | 62.645 | -3.682 | 1.00 17.53 | B_13 |
| MOTA | 2673 | CA | HIS | 129 | 47.001 | 63.366 | -4.718 | 1.00 26.87 | B_13 |
| MOTA | 2674 | CB | | | | | | | |
| | | | HIS | 129 | 47.495 | 62.398 | | | B_13 |
| MOTA | 2675 | CG | HIS | 129 | 48.729 | 61.645 | -5.400 | 1.00 19.64 | B_13 |
| MOTA | 2676 | CD2 | HIS | 129 | 49.769 | 61.996 | -4.609 | 1.00 19.96 | _ |
| | | _ | | | | | | | B_13 |
| MOTA | 2677 | NDI | HIS | 129 | 49.012 | 60.373 | -5.859 | 1.00 23.97 | B_13 |
| MOTA | 2679 | CE1 | HIS | 129 | 50.170 | 59.977 | -5.372 | 1.00 17.95 | B_13 |
| | | | | | | | | | • - |
| MOTA | 2680 | | HIS | 129 | 50.658 | 60.944 | -4.605 | 1.00 13.79 | B_13 |
| MOTA | 2681 | С | HIS | 129 | 46.153 | 64.457 | -5.360 | 1.00 39.97 | B_13 |
| MOTA | 2682 | 0 | HIS | 129 | 45.011 | | | | |
| | | | | | | 64.220 | -5.757 | 1.00 25.97 | B_13 |
| MOTA | 2683 | N | SER | 130 | 46.743 | 65.640 | -5.481 | 1.00 21.04 | B_13 |
| ATOM | 2685 | CA | SER | 130 | 46.090 | 66.776 | -6.109 | 1.00 16.72 | B_13 |
| | | | | | | _ | | | |
| MOTA | 2686 | CB | SER | 130 | 46.847 | 68.058 | - 5.757 | 1.00 20.97 | B_13 |
| MOTA | 2687 | QG | SER | 130 | 46.358 | 69.154 | -6.502 | 1.00 25.52 | B_13 |
| | | | | | | | | | |
| MOTA | 2689 | С | SER | 130 | 46.098 | 66.582 | -7.622 | 1.00 24.66 | B_13 |
| MOTA | 2690 | 0 | SER | 130 | 46.779 | 65.694 | -8.145 | 1.00 29.24 | B_13 |
| MOTA | 2691 | N | LYS | 131 | | | | | _ |
| | - | | | | 45.315 | 67.403 | -8.315 | 1.00 26.96 | B_13 |
| ATOM | 2693 | CA | LYS | 131 | 45.253 | 67.358 | -9.769 | 1.00 20.25 | B_13 |
| MOTA | 2694 | CB | LYS | 131 | 43.796 | 67 379 | -10.247 | 1.00 33.22 | |
| | | | | | _ | | | | B_13 |
| MOTA | 2695 | CG | LYS | 131 | 43.159 | 68.775 | -10.302 | 1.00 32.85 | B_13 |
| MOTA | 2696 | CD | LYS | 131 | 43.335 | 69 436 | -11.675 | 1.00 15.99 | B_13 |
| | | | | | | | | | |
| MOTA | 2697 | CE | LYS | 131 | 43.023 | 10.313 | -11.601 | 1.00 30.34 | B_13 |
| ATOM | 2698 | NZ | LYS | 131 | 43.879 | 71.647 | -10.600 | 1.00 30.44 | B_13 |
| MOTA | 2702 | С | LYS | 131 | 45.998 | | -10.249 | | _ |
| | | | | | | | | 1.00 15.31 | B_13 |
| MOTA | 2703 | 0 | LYS | 131 | 46.414 | 68.698 | -11.402 | 1.00 30.72 | B_13 |
| MOTA | 2704 | N | ASP | 132 | 46.191 | 69.536 | -9.323 | 1.00 23.41 | B_13 |
| | | | | • | | | | | |
| ATOM | 2706 | CA | ASP | 132 | 46.869 | 70.798 | | 1.00 22.69 | B_13 |
| MOTA | 2707 | CB | ASP | 132 | 46.641 | 71.726 | -8.379 | 1.00 24.86 | B_13 |
| MOTA | 2708 | CG | ASP | 132 | 46.819 | 73.200 | | 1.00 24.93 | B_13 |
| | | | | | | | | | |
| MOTA | 2709 | | ASP | | 46.007 | 74.009 | | 1.00 29.71 | B_13 |
| MOTA | 2710 | OD2 | ASP | 132 | 47.766 | 73.555 | -9.448 | 1.00 28.82 | B_13 |
| ATOM | 2711 | С | ASP | 132 | 48.358 | 70.497 | -9.728 | | |
| | | | • | | | | | 1.00 14.97 | B_13 |
| ATOM | 2712 | 0 | ASP | 132 | 49.047 | 70.235 | -8.742 | 1.00 19.64 | B_13 |
| ATOM | 2713 | N | PRO | 133 | 48.874 | | -10.964 | 1.00 16.94 | B_13 |
| | | | | | | | | | |
| MOTA | 2714 | CD | PRO | | 48.209 | | -12.199 | 1.00 21.42 | B_13 |
| ATOM | 2715 | CA | PRO | 133 | 50.293 | 70.264 | -11.215 | 1.00 19.34 | B_13 |
| | 2716 | | | | | | | | |
| MOTA | | CB | PRO | | 50.457 | | -12.690 | 1.00 20.48 | B_13 |
| MOTA | 2717 | CG | PRO | 133 | 49.347 | 71.636 | -12.929 | 1.00 21.80 | B_13 |
| ATOM | 2718 | C | PRO | | 51.237 | | -10.322 | | |
| | | | | | | | | 1.00 17.45 | B_13 |
| ATOM | 2719 | 0 | PRO | 133 | 52.319 | 70.590 | -10.006 | 1.00 23.30 | B_13 |
| MOTA | 2720 | N | GLY | | 50.799 | | | 1.00 32.46 | |
| | | | | | | | _ | | B_13 |
| MOTA | 2722 | CA | GLY | | 51.610 | 73.104 | | 1.00 19.44 | B_13 |
| MOTA | 2723 | C | GLY | 134 | 51.306 | 72.958 | | 1.00 22.33 | B_13 |
| ATOM | 2724 | | | | | | | | |
| | | 0 | GLY | | 51.556 | | | 1.00 21.92 | B_13 |
| MOTA | 2725 | N | ALA | 135 | 50.698 | 71.836 | -7.190 | 1.00 34.71 | B_13 |
| ATOM | 2727 | CA | ALA | | 50.355 | 71.580 | - | | D 13 |
| | | | | | | | | 1.00 18.35 | B_13 |
| ATOM | 2728 | CB | ALA | | 48.948 | 70.987 | -5.690 | 1.00 14.30 | B_13 |
| MOTA | 2729 | С | ALA | 135 | 51.370 | | - · · - · - | 1.00 10.00 | B_13 |
| ATOM | 2730 | ō | | | | | - · · - - | | |
| | | | ALA | | 51.739 | | · | 1.00 17.52 | B_13 |
| ATOM | 2731 | N | LEU | 136 | 51.727 | 70.842 | -3.952 | 1.00 21.29 | B_13 |
| | | | | | - 7 | | | | |
| | | | | | | | | | |

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| ATOM | 2733 | CA LEU | 136 | 52.692 | 70.015 | -3.230 | 1.00 14.62 | D 13 |
|------|------|---------|-----|--------|--------|---------|-------------|------|
| ATOM | 2734 | CB LEU | 136 | 52.738 | 70.458 | -1.763 | | B_13 |
| ATOM | 2735 | CG LEU | 136 | 54.007 | 70.308 | | 1.00 18.54 | B_13 |
| MOTA | 2736 | CD1 LEU | 136 | | | -0.921 | 1.00 34.11 | B_13 |
| MOTA | 2737 | | | 53.587 | 69.907 | 0.485 | 1.00 14.76. | B_13 |
| | | CD2 LEU | 136 | 54.969 | | -1.508 | 1.00 11.64 | B_13 |
| MOTA | 2738 | C LEU | 136 | 52.232 | 68.564 | -3.287 | 1.00 13.50 | B_13 |
| ATOM | 2739 | O LEU | 136 | 53.033 | 67.640 | -3.238 | 1.00 19.04 | B_13 |
| MOTA | 2740 | N MET | 137 | 50.921 | 68.364 | -3.281 | 1.00 17.54 | B_13 |
| MOTA | 2742 | CA MET | 137 | 50.360 | 67.019 | -3.324 | 1.00 25.11 | B_13 |
| ATOM | 2743 | CB MET | 137 | 49.010 | 66.981 | -2.599 | 1.00 19.80 | B_13 |
| MOTA | 2744 | CG MET | 137 | 49.083 | 67.312 | -1.117 | 1.00 15.35 | B_13 |
| MOTA | 2745 | SD MET | 137 | 50.354 | 66.361 | -0.262 | 1.00 11.22 | B_13 |
| ATOM | 2746 | CE MET | 137 | 49.882 | 64.680 | -0.764 | 1.00 13.90 | B_13 |
| ATOM | 2747 | C MET | 137 | 50.254 | 66.387 | -4.721 | 1.00 28.08 | B_13 |
| ATOM | 2748 | O MET | 137 | 49.730 | 65.268 | -4.863 | 1.00 12.18 | B_13 |
| ATOM | 2749 | N PHE | 138 | 50.771 | 67.070 | -5.743 | 1.00 10.00 | _ |
| ATOM | 2751 | CA PHE | 138 | 50.751 | 66.528 | -7.097 | 1.00 12.27 | B_13 |
| ATOM | 2752 | CB PHE | 138 | 51.327 | 67.523 | | 1.00 12.27 | B_13 |
| ATOM | 2753 | CG PHE | 138 | 51.051 | 67.175 | | • | B_13 |
| ATOM | 2754 | CD1 PHE | 138 | 52.090 | | | 1.00 25.74 | B_13 |
| ATOM | 2755 | CD2 PHE | 138 | | | -10.448 | 1.00 19.74 | B_13 |
| ATOM | 2756 | | | 49.747 | 67.007 | -9.990 | 1.00 24.46 | B_13 |
| ATOM | 2757 | CE1 PHE | 138 | 51.843 | | -11.786 | 1.00 19.54 | B_13 |
| | | CE2 PHE | 138 | 49.495 | | -11.335 | 1.00 24.12 | B_13 |
| ATOM | 2758 | CZ PHE | 138 | 50.544 | | -12.230 | 1.00 18.15 | B_13 |
| ATOM | 2759 | C PHE | 138 | 51.619 | 65.269 | -7.068 | 1.00 25.93 | B_13 |
| ATOM | 2760 | O PHE | 138 | 52.658 | 65.226 | | 1.00 12.50 | B_13 |
| MOTA | 2761 | N PRO | 139 | 51.166 | 64.194 | -7.714 | 1.00 25.17 | B_13 |
| MOTA | 2762 | CD PRO | 139 | 49.870 | 64.004 | -8.392 | 1.00 10.00 | B_13 |
| MOTA | 2763 | CA PRO | 139 | 51.950 | 62.956 | -7.713 | 1.00 18.48 | B_13 |
| MOTA | 2764 | CB PRO | 139 | 50.981 | 61.946 | -8.339 | 1.00 15.96 | B_13 |
| MOTA | 2765 | CG PRO | 139 | 50.140 | 62.798 | -9.250 | 1.00 18.82 | B_13 |
| MOTA | 2766 | C PRO | 139 | 53.299 | 62.950 | -8.430 | 1.00 17.22 | B_13 |
| MOTA | 2767 | O PRO | 139 | 53.849 | 61.876 | -8.661 | 1.00 36.93 | B_13 |
| MOTA | 2768 | N ILE | 140 | 53.844 | 64.114 | | 1.00 24.48 | B_13 |
| MOTA | 2770 | CA ILE | 140 | 55.118 | 64.155 | | 1.00 20.03 | B_13 |
| MOTA | 2771 | CB ILE | 140 | 54.996 | | -10.892 | 1.00 18.71 | |
| MOTA | 2772 | CG2 ILE | 140 | 56.334 | | -11.639 | 1.00 23.96 | B_13 |
| ATOM | 2773 | CG1 ILE | 140 | 53.932 | | -11.724 | | B_13 |
| ATOM | 2774 | CD1 ILE | 140 | 53.861 | | -13.125 | | B_13 |
| ATOM | 2775 | C ILE | 140 | 56.109 | | | 1.00 25.83 | B_13 |
| ATOM | 2776 | O ILE | | | 64.992 | -8.700 | 1.00 27.87 | B_13 |
| MOTA | 2777 | | 140 | 55.758 | 66.043 | -8.248 | 1.00 22.39 | B_13 |
| | | N TYR | 141 | 57.332 | 64.512 | -8.535 | 1.00 12.36 | B_13 |
| MOTA | 2779 | CA TYR | 141 | 58.350 | 65.281 | -7.834 | 1.00 21.85 | B_13 |
| MOTA | 2780 | CB TYR | 141 | 59.418 | 64.353 | -7.266 | 1.00 15.16 | B_13 |
| MOTA | 2781 | CG TYR | 141 | 60.592 | 65.096 | -6.672 | 1.00 15.65 | B_13 |
| ATOM | 2782 | CD1 TYR | 141 | 61.755 | 65.306 | -7.407 | 1.00 18.56 | B_13 |
| ATOM | 2783 | CE1 TYR | 141 | 62.836 | 65.967 | -6.859 | 1.00 10.00 | B_13 |
| ATOM | 2784 | CD2 TYR | 141 | 60.546 | 65.576 | -5.366 | 1.00 11.42 | B_13 |
| MOTA | 2785 | CE2 TYR | 141 | 61.626 | 66.236 | -4.814 | 1.00 13.45 | B_13 |
| ATOM | 2786 | CZ TYR | 141 | 62.770 | 66.429 | -5.567 | 1.00 10.00 | B_13 |
| MOTA | 2787 | OH TYR | 141 | 63.841 | 67.109 | -5.016 | 1.00 18.97 | B_13 |
| MOTA | 2789 | C TYR | 141 | 59.042 | 66.270 | -8.776 | 1.00 19:52 | B_13 |
| ATOM | 2790 | O ŢYR | 141 | 59.709 | 65.859 | -9.727 | 1.00 21.37 | B_13 |
| MOTA | 2791 | N THR | 142 | 58.932 | 67.556 | -8.465 | 1.00 23.99 | B_13 |
| MOTA | 2793 | CA THR | 142 | 59.573 | 68.616 | -9.238 | 1.00 19.53 | B_13 |
| MOTA | 2794 | CB THR | 142 | 58.515 | 69.578 | -9.807 | 1.00 10.00 | B_13 |
| MOTA | 2795 | OG1 THR | 142 | 57.704 | | -10.756 | 1.00 37.02 | B_13 |
| MOTA | 2797 | CG2 THR | 142 | 59.151 | | -10.457 | 1.00 34.35 | B_13 |
| MOTA | 2798 | C THR | 142 | 60.483 | 69.332 | -8.235 | 1.00 19.89 | B_13 |
| MOTA | 2799 | O THR | 142 | 60.120 | 69.513 | -7.076 | 1.00 25.67 | B_13 |
| MOTA | 2800 | N TYR | 143 | 61.699 | 69.677 | -8.643 | | B_13 |
| MOTA | 2802 | CA TYR | 143 | 62.609 | 70.344 | -7.707 | | B_13 |
| ATOM | 2803 | CB TYR | 143 | 64.091 | 70.190 | -8.108 | | |
| MOTA | 2804 | CG TYR | 143 | | | | 1.00 26.34 | B_13 |
| MOTA | 2805 | CD1 TYR | | 65.008 | 71.048 | -7.244 | | B_13 |
| MOTA | 2805 | | 143 | 65.066 | 70.866 | | | B_13 |
| | | CE1 TYR | 143 | 65.801 | 71.738 | -5.035 | 1.00 26.03 | B_13 |
| MOTA | 2807 | CD2 TYR | 143 | 65.714 | 72.114 | -7.795 | 1.00 17.36 | B_13 |
| ATOM | 2808 | CE2 TYR | 143 | 66.451 | 73.006 | | | B_13 |
| MOTA | 2809 | CZ TYR | 143 | 66.489 | 72.810 | | 1.00 10.00 | B_13 |
| MOTA | 2810 | OH TYR | 143 | 67.184 | 73.665 | | 1.00 27.84 | B_13 |
| MOTA | 2812 | C TYR | 143 | 62.330 | 71.815 | -7.456 | 1.00 24.77 | B_13 |
| MOTA | 2813 | O TYR | 143 | 62.201 | 72.611 | -8.399 | | B_13 |
| MOTA | 2814 | N THR | 144 | 62.292 | 72.160 | -6.170 | 1.00 22.23 | B_13 |
| MOTA | 2816 | CA THR | 144 | 62.103 | 73.533 | -5.727 | 1.00 33.68 | B_13 |
| ATOM | 2817 | CB THR | 144 | 60.668 | 73.814 | | | B_13 |
| MOTA | 2818 | OG1 THR | 144 | 60.277 | 72.812 | | 1.00 38.14 | B_13 |
| MOTA | 2820 | CG2 THR | 144 | 59.681 | 73.857 | -6.346 | 1.00 48.73 | B_13 |
| | | | | | | | ~ | |

| 37011 | 2021 | _ | | | 63 450 | | | | |
|-------|------|-----|------|-----|--------|--------|--------|------------|--------|
| ATOM | 2821 | C | THR | 144 | 63.178 | 73.893 | -4.695 | 1.00 35.52 | B_13 |
| ATOM | 2822 | 0 | THR | 144 | 64.207 | 74.465 | -5.064 | 1.00 39.57 | B_13 |
| MOTA | 2823 | N | GLY | 145 | 62.967 | 73.552 | -3.422 | 1.00 35.95 | B_13 |
| MOTA | 2825 | CA | GLY | 145 | 63.967 | 73.872 | -2.407 | 1.00 35.01 | B_13 |
| ATOM | 2826 | C | GLY | 145 | 63.509 | 74.025 | -0.965 | 1.00 26.81 | B_13 |
| MOTA | 2827 | 0 | GLY | 145 | 62.566 | 74.773 | -0.670 | 1.00 40.81 | B_13 |
| MOTA | 2828 | N | LYS | 146 | 64.302 | 73.439 | -0.066 | 1.00 27.13 | B_13 |
| MOTA | 2830 | CA | LYS | 146 | 64.071 | 73.423 | 1.389 | 1.00 23.89 | B_13 |
| ATOM | 2831 | CB | LYS | 146 | 65.163 | 72.548 | 2.049 | 1.00 29.08 | |
| ATOM | 2832 | CG | LYS | 146 | 64.992 | | | | B_13 |
| ATOM | 2833 | CD | LYS | | | 72.209 | 3.524 | 1.00 19.99 | B_13 |
| | | | | 146 | 66.079 | 71.224 | 3.913 | 1.00 20.44 | B_13 |
| ATOM | 2834 | CE | LYS | 146 | 66.181 | 71.010 | 5.402 | 1.00 24.16 | B_13 |
| MOTA | 2835 | NZ | LYS | 146 | 67.250 | 69.987 | 5.727 | 1.00 23.37 | B_13 |
| MOTA | 2839 | С | LYS | 146 | 63.926 | 74.778 | 2.124 | 1.00 18.98 | B_13 |
| MOTA | 2840 | 0 | LYS | 146 | 63.900 | 74.831 | 3.353 | 1.00 28.15 | B_13 |
| MOTA | 2841 | N | SER | 147 | 63.826 | 75.871 | 1.382 | 1.00 35.50 | B_13 |
| MOTA | 2843 | CA | SER | 147 | 63.661 | 77.185 | 1.992 | 1.00 31.59 | B_13 |
| ATOM | 2844 | CB | SER | 147 | 64.988 | 77.673 | 2.594 | 1.00 27.05 | B_13 |
| MOTA | 2845 | OG | SER | 147 | 65.996 | 77.756 | 1.586 | 1.00 48.28 | B_13 |
| MOTA | 2847 | C | SER | 147 | 63.203 | 78.131 | 0.902 | 1.00 27.12 | B_13 |
| MOTA | 2848 | 0 | SER | 147 | 62.743 | 79.251 | 1.168 | 1.00 33.75 | B_13 |
| ATOM | 2849 | N | HIS | 148 | 63.248 | 77.644 | -0.332 | 1.00 25.13 | |
| MOTA | 2851 | CA | HIS | 148 | 62.872 | 78.465 | | | B_13 |
| ATOM | 2852 | CB | HIS | 148 | | | -1.463 | 1.00 23.42 | B_13 |
| MOTA | 2853 | | | | 63.704 | 78.076 | -2.678 | 1.00 17.40 | B_13 |
| | | CG | HIS | 148 | 65.174 | 78.020 | -2.398 | 1.00 45.97 | B_13 |
| ATOM | 2854 | CD2 | | 148 | 66.204 | 77.524 | -3.121 | 1.00 27.24 | B_13 |
| MOTA | 2855 | | HIS | 148 | 65.724 | 78.476 | -1.213 | 1.00 43.49 | B_13 |
| MOTA | 2857 | | HIS | 148 | 67.024 | 78.253 | -1.218 | 1.00 30.28 | B_13 |
| MOTA | 2858 | NE2 | HIS | 148 | 67.342 | 77.676 | -2.366 | 1.00 45.28 | B_13 |
| MOTA | 2860 | С | HIS | 148 | 61.381 | 78.433 | -1.796 | 1.00 47.15 | B_13 |
| MOTA | 2861 | 0 | HIS | 148 | 60.936 | 79.166 | | _ | B_13 |
| ATOM | 2862 | N | PHE | 149 | 60.601 | 77.636 | | 1.00 48.76 | _ |
| ATOM | 2864 | CA | PHE | 149 | 59.170 | 77.557 | | | _ |
| MOTA | 2865 | CB | PHE. | 149 | 58.856 | 76.364 | | 1.00 27.77 | |
| ATOM | 2866 | CG | PHE | 149 | 58.415 | 76.781 | | | |
| ATOM | 2867 | | PHE | | | | | 1.00 24.63 | |
| | | | | 149 | 57.826 | 75.874 | - | 1.00 25.66 | B_13 |
| MOTA | 2868 | | PHE | 149 | 58.550 | 78.106 | | • | _ |
| MOTA | 2869 | | PHE | 149 | 57.376 | 76.277 | | | _ |
| MOTA | 2870 | | PHE | 149 | 58.104 | 78.520 | | | B_13 |
| MOTA | 2871 | CZ | PHE | 149 | 57.513 | 77.608 | -6.166 | 1.00 30.20 | B_13 |
| ATOM | 2872 | С | PHE | 149 | 58.061 | 77.791 | -0.308 | 1.00 27.40 | B_13 |
| ATOM | 2873 | 0 | PHE | 149 | 58.299 | 77.971 | | | |
| ATOM | 2874 | N | MET | 150 | 56.836 | 77.729 | | _ | B_13 |
| ATOM | 2876 | CA | MET | 150 | 55.621 | 78.027 | | | |
| MOTA | 2877 | CB | MET | 150 | 55.251 | 79.431 | _ | _ | _ |
| MOTA | 2878 | CG | MET | 150 | 55.599 | 79.691 | | _ | |
| ATOM | 2879 | SD | MET | 150 | 57.336 | | | | |
| ATOM | 2880 | CE | | | | | | 1.00 76.68 | _ |
| | | | MET | 150 | 57.209 | 81.473 | | | - |
| ATOM | 2881 | C | MET | 150 | 54.436 | 77.118 | _ | | |
| ATOM | 2882 | 0 | MET | 150 | 54.104 | | | | _ |
| MOTA | 2883 | N | LEU | 151 | 53.727 | 76.664 | | | B_13 |
| ATOM | 2885 | CA | LEU | 151 | 52.576 | 75.772 | _ | 1.00 25.68 | B_13 |
| MOTA | 2886 | CB | LEU | 151 | 51.968 | 75.474 | 1.807 | 1.00 23.46 | B_13 |
| ATOM | 2887 | CG | LEU | 151 | 51.087 | 74.232 | 1.927 | 1.00 24.21 | B_13 |
| MOTA | 2888 | CD1 | LEU | 151 | 51.936 | 72.998 | 1.657 | 1.00 21.54 | B_13 |
| ATOM | 2889 | CD2 | LEU | 151 | 50.487 | 74.150 | 3.314 | 1.00 19.89 | |
| MOTA | 2890 | С | LEU | 151 | 51.498 | 76.322 | | = | _ |
| ATOM | 2891 | 0 | LEU | 151 | 50.795 | 77.267 | | 1.00 35.38 | B_13 |
| ATOM | 2892 | N | PRO | 152 | 51.338 | 75.727 | -1.686 | 1.00 16.90 | B_13 |
| ATOM | 2893 | CD | PRO | 152 | 52.154 | 74.643 | -2.255 | 1.00 25.80 | |
| ATOM | 2894 | CA | PRO | 152 | 50.334 | 76.170 | | | B_13 |
| ATOM | 2895 | CB | | | | | -2.653 | 1.00 29.65 | B_13 |
| | | | PRO | 152 | 50.447 | 75.110 | -3.749 | 1.00 24.68 | B_13 |
| ATOM | 2896 | CG | PRO | 152 | 51.892 | 74.791 | -3.722 | 1.00 14.34 | B_13 |
| MOTA | 2897 | С | PRO | 152 | 48.910 | 76.261 | -2.087 | 1.00 10.00 | B_13 |
| MOTA | 2898 | 0 | PRO | 152 | 48.543 | 75.505 | -1.184 | 1.00 20.25 | B_13 |
| ATOM | 2899 | N | ASP | 153 | 48.117 | 77.180 | | 1.00 19.53 | B_13 |
| MOTA | 2901 | CA | ASP | 153 | 46.723 | 77.387 | -2.226 | 1.00 15.90 | B_13 |
| ATOM | 2902 | CB | ASP | 153 | 45.986 | 78.304 | -3.213 | 1.00 22.34 | B_13 |
| ATOM | 2903 | CG | ASP | 153 | 46.418 | 79.741 | -3.115 | 1.00 28.86 | B_13 |
| ATOM | 2904 | | ASP | 153 | 47.016 | 80.115 | -2.074 | 1.00 25.86 | |
| ATOM | 2905 | | ASP | 153 | 46.142 | 80.494 | | | B_13 |
| ATOM | 2906 | C | ASP | 153 | 45.953 | | -4.084 | 1.00 30.09 | B_13 |
| MOTA | 2907 | Ö | | | | 76.084 | -2.169 | | B_13 |
| ATOM | 2908 | | ASP | 153 | 45.309 | 75.783 | -1.167 | | B_13 |
| ATOM | | N | ASP | 154 | 46.000 | 75.339 | -3.276 | 1.00 25.51 | B_13 |
| | 2910 | CA | ASP | 154 | 45.316 | 74.063 | -3.392 | 1.00 20.91 | B_13 |
| MOTA | 2911 | CB | ASP | 154 | 45.745 | 73.364 | -4.682 | 1.00 14.23 | · B_13 |
| ATOM | 2912 | CG | ASP | 154 | 45.033 | 72.062 | -4.885 | 1.00 22.95 | B_13 |
| | | | | | | | | | |

| ATOM | 2913 | OD1 | ASP | 154 | 45.590 | 71.026 | -4.516 | 1.00 17.80 | B_13 |
|--------------|--------------|------------|------------|------------|------------------|------------------|------------------|--------------------------|--------------|
| MOTA | 2914 | | ASP | 154 | 43.904 | 72.076 | -5.388 | 1.00 19.14 | B_13 |
| MOTA | 2915 | | ASP | 154 | 45.551 | 73.155 | -2.173 | 1.00 26.95 | B_13 |
| ATOM | 2916 | | ASP | 154 | 44.629 | 72.491 | -1.696 | 1.00 22.92 1.00 23.56 | B_13 |
| MOTA | 2917 2919 | | ASP ASP | 155 155 | 46.776 47.110 | 73.155 72.338 | -1.654 -0.490 | 1.00 23.56 1.00 28.69 | B_13 B_13 |
| MOTA MOTA | 2920 | | ASP | 155 | 48.618 | 72.118 | -0.388 | 1.00 12.87 | B_13 |
| ATOM | 2921 | • | ASP | 155 | 49.208 | 71.566 | -1.676 | 1.00 24.35 | B_13 |
| MOTA | 2922 | | ASP | 155 | 49.705 | 72.369 | -2.500 | 1.00 27.89 | B_13 |
| ATOM | 2923 | | ASP | 155 | 49.152 | 70.335 | -1.875 | 1.00 16.96 | B_13 |
| ATOM | 2924 | | ASP | 155 | 46.582 | 72.976 | 0.781 | 1.00 25.41 1.00 13.36 | B_13 |
| ATOM | 2925 2926 | | ASP VAL | 155 156 | 46.055 46.733 | 72.275 74.296 | 1.656 0.891 | 1.00 16.99 | B_13 B_13 |
| MOTA ATOM | 2928 | | VAL | 156 | 46.222 | 75.021 | 2.053 | 1.00 22.26 | B_13 |
| ATOM | 2929 | | VAL | 156 | 46.340 | 76.571 | 1.901 | 1.00 25.69 | B_13 |
| MOTA | 2930 | | VAL | 156 | 45.811 | 77.249 | 3.158 | 1.00 14.95 | B_13 |
| MOTA | 2931 | | VAL | 156 | 47.768 | 77.007 | 1.641 | 1.00 17.52 | B_13 |
| MOTA | 2932 | | VAL | 156 156 | 44.727 | 74.705 74.234 | 2.129 3.145 | 1.00 10.00 1.00 22.47 | B_13 B_13 |
| MOTA MOTA | 2933 2934 | O N | VAL GLN | 156 157 | 44.224 44.033 | 74.234 | 1.029 | 1.00 22.47 | B_13 |
| ATOM | 2936 | CA | GLN | 157 | 42.604 | 74.758 | 0.930 | 1.00 17.97 | B_13 |
| ATOM | 2937 | CB | GLN | 157 | 42.108 | 75.039 | -0.497 | 1.00 17.10 | B_13 |
| MOTA | 2938 | CG | GLN | 157 | 40.804 | 75.852 | -0.547 | 1.00 26.00 | B_13 |
| MOTA | 2939 | CD | GLN | 157 | 40.949 | 77.284 | -0.005 | 1.00 25.84 1.00 39.61 | B_13 |
| ATOM | 2940 2941 | OE1 NE2 | GLN GLN | 157 157 | 41.218 40.744 | 77.505 78.255 | 1.177 -0.875 | 1.00 39.61 1.00 32.22 | B_13 B_13 |
| MOTA MOTA | 2944 | C | GLN | 157 | 42.347 | 73.324 | 1.309 | 1.00 18.69 | B_13 |
| MOTA | 2945 | Ö | GLN | 157 | 41.368 | 73.015 | 1.982 | 1.00 10.00 | B_13 |
| ATOM | 2946 | N | GLY | 158 | 43.272 | 72.460 | 0.903 | 1.00 31.05 | B_13 |
| MOTA | 2948 | CA | GLY | 158 | 43.156 | 71.053 | 1.205 | 1.00 21.69 | B_13 |
| ATOM | 2949 | C | GLY | 158 | 43.129 | 70.738 | 2.684 | 1.00 13.51 1.00 14.91 | B_13 |
| MOTA MOTA | 2950 2951 | И О | GLY ILE | 158 159 | 42.108 44.224 | 70.263 71.006 | 3.182 3.398 | 1.00 14.91 | B_13 B_13 |
| MOTA | 2953 | CA | ILE | 159 | 44.268 | 70.686 | 4.827 | 1.00 19.14 | B_13 |
| MOTA | 2954 | CB | ILE | 159 | 45.669 | 70.880 | 5.503 | 1.00 12.57 | B_13 |
| MOTA | 2955 | CG2 | ILE | 159 | 46.268 | 69.542 | 5.960 | 1.00 19.22 | B_13 |
| MOTA | 2956 | CG1 | | 159 | 46.603 | 71.702 | 4.633 | 1.00 31.62 | B_13 |
| MOTA | 2957 | CD1 | ILE | 159 150 | 46.426 | 73.177 | 4.824 5.610 | 1.00 25.87 1.00 21.87 | B_13 B_13 |
| MOTA MOTA | 2958 2959 | C O | ILE | 159 159 | 43.235 42.691 | 71.461 70.952 | 6.592 | 1.00 21.07 | B_13 |
| MOTA | 2960 | N | GLN | 160 | 42.959 | 72.689 | 5.186 | 1.00 12.08 | B_13 |
| ATOM | 2962 | CA | GLN | 160 | 41.967 | 73.483 | 5.874 | 1.00 11.43 | B_13 |
| MOTA | 2963 | CB | GLN | 160 | 41.949 | 74.916 | 5.346 | 1.00 29.25 | B_13 |
| MOTA | 2964 | CG | GLN | 160 | 43.158 | 75.737 | 5.827 | 1.00 22.01 | B_13 |
| ATOM | 2965 2966 | CD OE1 | GLN GLN | 160 160 | 43.098 42.260 | 77.199 77.593 | 5.416 4.607 | 1.00 18.77 1.00 36.02 | B_13 B_13 |
| MOTA MOTA | 2967 | NE2 | | 160 | 43.997 | 78.004 | 5.965 | 1.00 28.49 | B_13 |
| ATOM | 2970 | C | GLN | 160 | 40.596 | | | 1.00 22.28 | B_13 |
| MOTA | 2971 | 0 | GLN | 160 | 39.855 | | | 1.00 14.16 | B_13 |
| ATOM | 2972 | N | SER | 161 | 40.304 | 72.183 | | 1.00 32.89 | B_13 |
| ATOM | 2974 2975 | CA CB | SER SER | 161 161 | 39.005 38.847 | 71.537 70.901 | | | B_13 B_13 |
| MOTA MOTA | 2976 | OG | SER | 161 | 39.594 | | | 1.00 24.88 | B_13 |
| MOTA | 2978 | C | SER | 161 | 38.831 | | | | B_13 |
| MOTA | 2979 | 0 | SER | 161 | 37.745 | | | | B_13 |
| ATOM | 2980 | N | LEU | 162 | 39.931 | 69.852 | | | B_13 |
| MOTA MOTA | 2982 2983 | CA CB | LEU LEU | 162 162 | 39.913 41.081 | 68.829 67.852 | | 1.00 29.17 1.00 12.08 | B_13 B_13 |
| MOTA | 2984 | | LEU | 162 | 40.982 | | | | B_13 |
| ATOM | 2985 | | | 162 | 40.661 | | | | B_13 |
| MOTA | 2986 | | | 162 | 42.299 | | | | B_13 |
| MOTA | 2987 | | LEU | 162 | 39.965 | | | | B_13 |
| MOTA | 2988 | | LEU | 162 163 | 39.047 41.015 | | | | B_13 B_13 |
| MOTA MOTA | 2989 2991 | | TYR TYR | 163 | 41.211 | | | | B_13 |
| ATOM | 2992 | | TYR | 163 | 42.695 | | | - | B_13 |
| MOTA | 2993 | | TYR | 163 | 43.221 | 69.167 | 10.209 | 1.00 10.00 | B_13 |
| MOTA | 2994 | | | 163 | 43.114 | | | | B_13 |
| MOTA | 2995 | | TYR | 163 | 43.452 | | | | B_13 |
| MOTA MOTA | 2996 2997 | | TYR TYR | 163 163 | 43.703 44.048 | | | | B_13 B_13 |
| MOTA | 2998 | | TYR | 163 | 43.914 | | | | B_13 |
| MOTA | 2999 | | TYR | 163 | 44.210 | 65.121 | 9.711 | | B_13 |
| MOTA | 3001 | | TYR | 163 | 40.634 | 72.085 | 10.187 | | B_13 |
| MOTA | 3002 | | TYR | 163 | 39.975 | | | | B_13 |
| MOTA MOTA | 3003 3005 | | GLY GLY | 164 164 | 40.819 40.291 | | | | B_13 B_13 |
| 414 VII | -000 | ~~ | ani | 704 | TV . 431 | | . 5.340 | 2.00 30.04 | 2_10 |

| | 2006 | | 1.64 | 44 400 | | | | |
|------|------|---------|-------|--------|--------|-----------------|------------|--------|
| MOTA | 3006 | C GLY | 164 | 41.402 | 75.344 | 9.424 | 1.00 30.89 | B_13 |
| ATOM | 3007 | O GLY | 164 | 41.101 | 76.564 | 9.368 | 1.00 26.89 | B_13 |
| MOTA | 3008 | OT GLY | 164 | 42.570 | 74.911 | 9.560 | 1.00 27.71 | B_13 |
| MOTA | 3013 | ZN ZN | 166 | 51.961 | 60.891 | -2.865 | 1.00 28.31 | BION |
| ATOM | 3014 | ZN ZN | 167 | 56.468 | 50.981 | 3.458 | 1.00 26.20 | |
| | | | | | | | | BION |
| MOTA | 3015 | | 168 | 63.096 | 53.752 | -5.445 | 1.00 14.89 | BION |
| MOTA | 3016 | CA CA | 165 | 50.705 | 55.618 | 13.085 | 1.00 15.79 | BION |
| MOTA | 3047 | C5 WAY | 169 | 54.585 | 56.119 | -6.288 | 1.00 40.09 | B693 |
| MOTA | 3048 | CF1 WAY | 169 | 54.019 | 54.934 | -5.802 | 1.00 21.52 | B693 |
| | | | | | | | | |
| MOTA | 3049 | CH WAY | 169 | 53.271 | 54.923 | -4.624 | 1.00 32.32 | B693 |
| MOTA | 3050 | C2 WAY | 169 | 53.100 | 56.104 | -3.898 | 1.00 21.39 | B693 |
| MOTA | 3051 | C3 WAY | 169 | 53.667 | 57.286 | -4.369 | 1.00 18.26 | B693 |
| MOTA | 3052 | C4 WAY | 169 | 54.402 | 57.308 | -5.540 | 1.00 20.63 | B693 |
| MOTA | 3053 | N20 WAY | | 54.933 | 58.531 | -5.964 | 1.00 22.15 | B693 |
| | | | | | | | | |
| MOTA | 3054 | CD WAY | | 54.297 | 59.340 | -7.031 | 1.00 30.92 | B693 |
| MOTA | 3055 | C23 WAY | 169 | 53.576 | 58.491 | -8.087 | 1.00 20.75 | B693 |
| MOTA | 3056 | C28 WAY | 169 | 54.224 | 58.114 | -9.279 | 1.00 34.14 | B693 |
| MOTA | 3057 | C27 WAY | 169 | 53.539 | 57.335 | -10.228 | 1.00 33.99 | B693 |
| MOTA | 3058 | CM WAY | | 52.209 | 56.944 | -9.968 | 1.00 23.49 | B693 |
| | | | | | | | • | |
| MOTA | 3059 | N25 WAY | | 51.602 | 57.318 | -8.814 | 1.00 23.61 | B693 |
| MOTA | 3060 | C24 WAY | | 52.246 | 58.071 | -7.880 | 1.00 20.52 | B693 |
| MOTA | 3061 | S21 WAY | 169 | 56.531 | 58.783 | -5.660 | 1.00 20.46 | B693 |
| MOTA | 3062 | C16 WAY | 169 | 56.457 | 60.446 | -5.010 | 1.00 39.00 | B693 |
| MOTA | 3063 | C21 WAY | | 56.700 | 60.669 | -3.634 | 1.00 28.79 | B693 |
| | 3064 | | | | | | | |
| ATOM | | C20 WAY | | 56.656 | 61.967 | -3.109 | 1.00 12.65 | B693 |
| MOTA | 3065 | C19 WAY | | 56.373 | 63.058 | -3.946 | 1.00 15.68 | B693 |
| ATOM | 3066 | C18 WAY | 169 | 56.126 | 62.828 | -5.319 | 1.00 12.08 | B693 |
| ATOM | 3067 | C17 WAY | 169 | 56.169 | 61.538 | -5.852 | 1.00 15.19 | B693 |
| ATOM | 3068 | O33 WAY | | 56.337 | 64.360 | -3.424 | 1.00 16.79 | B693 |
| | 3069 | | | | | | | |
| MOTA | | C36 WAY | | 56.982 | 65.456 | -4.084 | 1.00 20.80 | B693 |
| ATOM | 3070 | O15 WAY | | 56.973 | 57.923 | -4.580 | 1.00 21.90 | B693 . |
| ATOM | 3071 | O14 WAY | 169 | 57.259 | 58.799 | -6.913 | 1.00 10.86 | B693 |
| MOTA | 3072 | C7 WAY | 169 | 53.486 | 58.556 | -3.613 | 1.00 10.00 | B693 |
| ATOM | 3073 | N9 WAY | | 53.741 | 58.606 | -2.303 | 1.00 10.00 | B693 |
| | 3074 | | | | | | | |
| ATOM | | O10 WAY | | 53.539 | 59.846 | -1.659 | 1.00 23.73 | B693 |
| ATOM | 3075 | OS WAY | | 53.107 | 59.569 | -4.154 | 1.00 15.89 | B693 |
| MOTA | 3076 | C29 WAY | 169 | 55.383 | 55.968 | -7.606 | 1.00 28.30 | B693 |
| ATOM | 1 | OH2 WAT | 301 | 67.399 | 53.332 | 19.612 | 1.00 10.00 | SOLV |
| ATOM | 2 | OH2 WAT | | 61.288 | 46.506 | 17.898 | 1.00 10.00 | SOLV |
| ATOM | 3 | OH2 WAT | | | | | | |
| | _ | | | 79.538 | 50.433 | 20.115 | 1.00 10.00 | SOLV |
| ATOM | 4 | OH2 WAT | | 80.982 | 25.236 | 19.076 | 1.00 26.37 | SOLV |
| ATOM | 5 | OH2 WAT | 305 | 82.461 | 30.767 | 19.346 | 1.00 13.02 | SOLV |
| ATOM | 6 | OH2 WAT | 306 | 67.759 | 41.912 | 4.887 | 1.00 17.30 | SOLV |
| ATOM | 7 | OH2 WAT | | 60.785 | 41.727 | 10.585 | 1.00 20.42 | SOLV |
| ATOM | 8 | OH2 WAT | | 89.638 | 33.523 | | | |
| | | | | | | 25.640 | 1.00 33.45 | SOLV |
| ATOM | 9 | OH2 WAT | | 77.721 | 51.975 | 4.391 | 1.00 13.91 | |
| MOTA | 10 | OH2 WAT | _ | 96.022 | 34.702 | 6.692 | 1.00 25.50 | SOLV |
| ATOM | 11 | OH2 WAI | 311 | 71.292 | 38.746 | 26.741 | 1.00 13.06 | SOLV |
| MOTA | 12 | OH2 WAT | 312 | 85.939 | 49.781 | 3.498 | 1.00 12.04 | SOLV |
| MOTA | 13 | OH2 WAT | | 58.101 | 41.127 | 10.261 | 1.00 40.97 | SOLV |
| ATOM | 14 | OH2 WAT | | 86.373 | 42.692 | 0.747 | 1.00 17.24 | |
| | | | | | | | | SOLV |
| MOTA | 15 | OH2 WAT | | 78.257 | 39.885 | 24.626 | 1.00 18.57 | SOLV |
| MOTA | 16 | OH2 WAT | | 68.341 | 48.572 | 25.558 | 1.00 18.33 | SOLV |
| MOTA | 17 | OH2 WAT | 317 | 79.806 | 29.147 | 18.371 | 1.00 10.00 | SOLV |
| ATOM | 18 | OH2 WAT | 318 | 87.119 | 44.480 | 23.137 | 1.00 46.31 | SOLV |
| MOTA | 19 | OH2 WAT | | 55.885 | 39.688 | 11.459 | 1.00 21.26 | SOLV |
| ATOM | 20 | | | | | | | |
| | | | | 73.250 | 41.084 | 0.386 | 1.00 18.49 | SOLV |
| ATOM | 21 | OH2 WAT | | 72.079 | 46.488 | -6.835 | 1.00 27.48 | SOLV |
| MOTA | 22 | OH2 WAT | 322 | 71.923 | 37.638 | ~3.750 | 1.00 29.19 | SOLV |
| MOTA | 23 | OH2 WAT | 323 | 74.998 | 28.451 | 2.684 | 1.00 34.60 | SOLV |
| ATOM | 24 | OH2 WAT | | 87.769 | 44.123 | 9.214 | 1.00 15.60 | SOLV |
| MOTA | 25 | | | 86.113 | | | | |
| | | | | | 24.382 | 16.709 | 1.00 25.17 | SOLV |
| MOTA | 26 | | | 81.205 | 57.603 | | 1.00 34.27 | SOLV |
| MOTA | 27 | | | 75.163 | 62.739 | 12.391 | 1.00 16.47 | SOLV |
| MOTA | 28 | OH2 WAT | 328 | 65.604 | 44.690 | 2.830 | 1.00 26.64 | SOLV |
| ATOM | 29 | | | 61.899 | 45.512 | | 1.00 15.82 | SOLV |
| MOTA | . 30 | | | 58.763 | 41.730 | | | |
| | | | | | | - | 1.00 27.95 | SOLV |
| MOTA | 31 | | | 69.823 | 44.729 | | 1.00 13.37 | SOLV |
| MOTA | 32 | | | 79.220 | 61.263 | 12.781 | 1.00 28.84 | SOLV |
| MOTA | 33 | OH2 WAT | r 333 | 78.105 | 37.095 | 27.911 | 1.00 34.48 | SOLV |
| MOTA | 34 | | | 75.939 | | | 1.00 35.21 | SOLV |
| MOTA | 35 | | | 90.256 | 42.668 | 16.539 | 1.00 45.05 | SOLV |
| ATOM | 36 | | | | | | | |
| | | | | 86.761 | 51.457 | 13.881 | 1.00 25.26 | SOLV |
| ATOM | 37 | | | 67.479 | | -5.009 | 1.00 33.30 | SOLV |
| MOTA | 38 | | | 82.018 | 50.963 | 8.823 | 1.00 19.80 | SOLV |
| MOTA | 39 | | r 339 | 80.278 | 32.895 | -1.126 | 1.00 30.16 | SOLV |
| MOTA | 40 | OH2 WAT | r 340 | 71.683 | 50.944 | 31.567 | 1.00 29.62 | SOLV , |
| | | | | | | - - | | |
| | | | | | | | | |

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| MOTA | 41 | OH2 WAT | 341 | 61.633 | 49.360 | 10.951 | 1.00 15.47 | SOLV |
|--------|------|---------|-----|--------|--------|---------|------------|------|
| MOTA | 42 | OH2 WAT | 342 | 89.589 | 43.811 | 5.959 | 1.00 18.08 | SOLV |
| MOTA | 43 | OH2 WAT | 343 | 70.742 | 35.952 | 14.932 | 1.00 34.03 | SOLV |
| MOTA | 44 | OH2 WAT | 344 | 89.836 | 28.590 | 26.657 | 1.00 18.11 | SOLV |
| ATOM | 45 | OH2 WAT | 345 | 70.822 | 32.764 | 1.461 | 1.00 22.35 | SOLV |
| MOTA | 46 | OH2 WAT | 346 | 63.056 | 34.653 | 0.491 | 1.00 29.51 | |
| ATOM | 47 | OH2 WAT | 347 | 58.054 | 46.282 | | | SOLV |
| | | | | | | 2.363 | 1.00 10.00 | SOLV |
| MOTA | 48 | OH2 WAT | 348 | 67.914 | 58.660 | -6.267 | 1.00 18.30 | SOLV |
| MOTA | · 49 | OH2 WAT | 349 | 70.170 | 56.725 | 0.575 | 1.00 11.89 | SOLV |
| MOTA | 50 | OH2 WAT | 350 | 55.922 | 73.897 | 0.623 | 1.00 18.86 | SOLV |
| ATOM | 51 | OH2 WAT | 351 | 73.489 | 53.195 | 2.061 | 1.00 24.35 | SOLV |
| ATOM | 52 | OH2 WAT | 352 | 58.033 | 50.530 | 19.075 | 1.00 25.52 | SOLV |
| MOTA | 53 | OH2 WAT | 353 | 63.245 | 57.302 | 17.340 | 1.00 13.88 | SOLV |
| MOTA | 54 | OH2 WAT | 354 | 58.442 | 71.334 | -5.670 | 1.00 17.51 | SOLV |
| ATOM | 55 | OH2 WAT | 355 | 62.535 | 61.154 | 16.706 | 1.00 12.38 | SOLV |
| ATOM | 56 | OH2 WAT | 356 | 66.949 | 51.163 | | 1.00 17.92 | SOLV |
| MOTA | 57 | OH2 WAT | 357 | 57.588 | 54.191 | 9.850 | 1.00 17.88 | SOLV |
| MOTA | 58 | OH2 WAT | 358 | 64.836 | 48.085 | 4.627 | 1.00 17.80 | SOLV |
| MOTA | 59 | OH2 WAT | 359 | 66.445 | 61.785 | 19.640 | 1.00 24.12 | SOLV |
| ATOM | 60 | OH2 WAT | 360 | 55.740 | 42.557 | 0.533 | 1.00 27.32 | |
| ATOM | 61 | OH2 WAT | 361 | 74.075 | 57.146 | 13.179 | | SOLV |
| | 62 | OH2 WAT | 362 | | | | | SOLV |
| MOTA | | | | 46.987 | 69.315 | -2.545 | 1.00 11.87 | SOLV |
| ATOM | 63 | OH2 WAT | 363 | 53.842 | 52.266 | -2.612 | 1.00 25.20 | SOLV |
| ATOM | 64 | OH2 WAT | 364 | 33.425 | 65.313 | -4.686 | 1.00 28.97 | SOLV |
| MOTA | 65 | CH2 WAT | 365 | 45.633 | 51.173 | 10.502 | 1.00 31.97 | SOLV |
| MOTA | 66 | OH2 WAT | 366 | 39.040 | 71.050 | -0.722 | 1.00 20.81 | SOLV |
| MOTA | 67 | OH2 WAT | 367 | 54.517 | 67.335 | -6.251 | 1.00 46.24 | SOLV |
| MOTA | 68 | OH2 WAT | 368 | 45.083 | 67.138 | 20.314 | 1.00 29.47 | SOLV |
| ATOM | 69 | OH2 WAT | 369 | 65.758 | 67.669 | -6.655 | 1.00 14.69 | SOLV |
| ATOM | 70 | OH2 WAT | 370 | 44.943 | 78.174 | 12.948 | 1.00 23.88 | SOLV |
| MOTA | 71 | OH2 WAT | 371 | 37.141 | 57.403 | 1.723 | 1.00 23.72 | SOLV |
| ATOM | 72 | OH2 WAT | 372 | 62.407 | 66.806 | 13.368 | | SOLV |
| MOTA | 73 | OH2 WAT | 373 | 50.776 | 47.263 | 5.661 | 1.00 38.22 | SOLV |
| ATOM | 74 | OH2 WAT | 374 | 56.697 | 47.264 | 11.752 | 1.00 24.75 | SOLV |
| ATOM | 75 | OH2 WAT | 375 | 42.566 | 60.884 | 15.739 | 1.00 24.75 | |
| ATOM | 76 | OH2 WAT | | | | | | SOLV |
| | | | 376 | 59.299 | | 13.838 | 1.00 31.27 | SOLV |
| MOTA | 77 | OH2 WAT | 377 | 72.976 | 63.691 | -0.667 | 1.00 20.36 | SOLV |
| MOTA | 78 | OH2 WAT | 378 | 72.876 | 60.516 | -6.752 | 1.00 34.24 | SOLV |
| MOTA | 79 | OH2 WAT | 379 | 63.998 | 68.760 | 16.371 | | SOLV |
| MOTA | 80 | OH2 WAT | 380 | 44.947 | 66.728 | -2.566 | 1.00 29.51 | SOLV |
| MOTA | 81 | OH2 WAT | 381 | 57.690 | 61.926 | -9.414 | 1.00 29.01 | SOLV |
| MOTA | 82 | OH2 WAT | 382 | 44.595 | 80.810 | 5.831 | 1.00 27.43 | SOLV |
| ATOM | 83 | OH2 WAT | 383 | 78.065 | 36.583 | 24.121 | 1.00 14.08 | SOLV |
| ATOM . | 84 | OH2 WAT | 384 | 42.289 | 64.651 | -0.868 | 1.00 25.57 | SOLV |
| MOTA | 85 | OH2 WAT | 385 | 59.851 | 68.458 | ~12.381 | 1.00 30.18 | SOLV |
| ATOM | 86 | OH2 WAT | 386 | 53.784 | | | 1.00 22.35 | SOLV |
| MOTA | - 87 | OH2 WAT | 387 | 72.793 | | 8.925 | | SOLV |
| MOTA | 88 | OH2 WAT | 388 | 57.224 | 68.062 | -6.072 | 1.00 17.87 | SOLV |
| MOTA | 89 | OH2 WAT | 389 | 45.210 | 44.988 | 4.285 | 1.00 25.10 | SOLV |
| ATOM | 90 | OH2 WAT | 390 | 49.413 | 53.782 | 1.546 | | SOLV |
| ATOM | 91 | OH2 WAT | 391 | 45.232 | | 1.393 | 1.00 19.25 | |
| ATOM | 92 | OH2 WAT | 392 | 42.551 | | | | SOLV |
| ATOM | 93 | OH2 WAT | | | 59.954 | 5.056 | 1.00 27.30 | SOLV |
| | | | 393 | 58.412 | 43.750 | 3.948 | 1.00 58.70 | SOLV |
| ATOM | 94 | OH2 WAT | 394 | 56.942 | 54.199 | -2.588 | 1.00 31.14 | SOLV |
| ATOM | 95 | OH2 WAT | 395 | 55.216 | 51.994 | 9.824 | 1.00 13.25 | SOLV |
| MOTA | 96 | OH2 WAT | 396 | 51.642 | | 14.874 | 1.00 10.00 | SOLV |
| ATOM | 97 | OH2 WAT | 397 | 48.690 | | 13.991 | 1.00 28.59 | SOLV |
| MOTA | 98 | OH2 WAT | 398 | 74.412 | 37.913 | 0.396 | 1.00 12.55 | SOLV |
| MOTA | 99 | OH2 WAT | 399 | 81.920 | 53.968 | 18.267 | 1.00 14.05 | SOLV |
| MOTA | 100 | OH2 WAT | 400 | 70.413 | 41.780 | 1.170 | 1.00 16.68 | SOLV |
| MOTA | 101 | OH2 WAT | 401 | 71.098 | 53.544 | 2.407 | 1.00 27.63 | SOLV |
| MOTA | 102 | OH2 WAT | 402 | 94.383 | 32.979 | 9.497 | 1.00 27.97 | SOLV |
| MOTA | 103 | OH2 WAT | 403 | 70.765 | 66.069 | 16.389 | 1.00 38.09 | SOLV |
| ATOM | 104 | OH2 WAT | 404 | 78.651 | 34.890 | 29.495 | 1.00 48.60 | SOLV |
| ATOM | 105 | OH2 WAT | 405 | 80.289 | 39.811 | 24.727 | 1.00 20.74 | SOLV |
| ATOM | 106 | OH2 WAT | 406 | 63.627 | 47.414 | 7.301 | 1.00 20.74 | SOLV |
| MOTA | 107 | OH2 WAT | 407 | 74.679 | 30.772 | | | |
| MOTA | 108 | OH2 WAT | 408 | | | 11.524 | 1.00 37.03 | SOLV |
| ATOM | 109 | | | 80.240 | 36.041 | 26.681 | 1.00 27.42 | SOLV |
| | | OH2 WAT | 409 | 84.971 | 25.909 | 18.426 | 1.00 24.96 | SOLV |
| MOTA | 110 | OH2 WAT | 410 | 57.832 | 41.294 | 5.792 | 1.00 71.90 | SOLV |
| MOTA | 111 | OH2 WAT | 411 | 55.484 | | -9.086 | 1.00 48.47 | SOLV |
| MOTA | 112 | OH2 WAT | 412 | 65.535 | 68.260 | 2.400 | 1.00 26.24 | SOLV |
| ATOM | 113 | OH2 WAT | 413 | 80.085 | 42.291 | -3.144 | 1.00 26.49 | SOLV |
| MOTA | 114 | OH2 WAT | 414 | 82.088 | 37.456 | | 1.00 42.54 | SOLV |
| ATOM | 115 | OH2 WAT | 415 | 61.020 | 53.195 | 21.566 | 1.00 38.16 | SOLV |
| MOTA | 116 | OH2 WAT | 416 | 55.968 | 70.365 | -5.096 | 1.00 28.42 | SOLV |
| MOTA | 117 | OH2 WAT | 417 | 51.619 | 57.620 | -0.487 | 1.00 41.81 | SOLV |

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| MOTA | 118 | OH2 WAT | 418 | 40.651 | 66.108 | 2.086 | 1.00 40.11 | SOLV |
|------|-----|---------|-----|----------|--------|--------|------------|------|
| ATOM | 119 | OH2 WAT | 419 | · 58.453 | 49.818 | 7.926 | 1.00 38.96 | SOLV |
| MOTA | 120 | OH2 WAT | 420 | 53.768 | 51.716 | 13.623 | 1.00 43.62 | SOLV |
| MOTA | 121 | OH2 WAT | 421 | 76.068 | 60.373 | 21.292 | 1.00 39.30 | SOLV |
| ATOM | 122 | OH2 WAT | 422 | 56.186 | 50.034 | 17.422 | 1.00 37.47 | SOLV |
| END | | | | | | | | |

FIG. 6

Compound C

FIG. 7

75/75



SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/05150

| A. *CLASSIFICATION OF SUBJECT MATTER | | | | | | | | |
|---|--|---|-----------------------|--|--|--|--|--|
| IPC(7) :G01N 9/00, 33/48 US CL :435/183; 702/22 | | | | | | | | |
| According to International Patent Classification (IPC) or to both national classification and IPC | | | | | | | | |
| B. FIELDS SEARCHED | | | | | | | | |
| Minimum d | ocumentation searched (classification system followed | by classification symbols) | ļ | | | | | |
| U.S. : 435/183; 702/22 | | | | | | | | |
| Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched | | | | | | | | |
| NONE . | | | | | | | | |
| Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) | | | | | | | | |
| STN: WE | STN: WEST | | | | | | | |
| | | | | | | | | |
| C. DOCUMENTS CONSIDERED TO BE RELEVANT | | | | | | | | |
| Category* | Citation of document, with indication, where ap | propriate, of the relevant passages | Relevant to claim No. | | | | | |
| X | GOMIS-RUTH, F.X. et al. The he | | 8-14 | | | | | |
| | (MMP-13: 2.7, ANG> crystal st haemopexin-like domain. Journal Mol | | | | | | | |
| | 3, pages 556-566, see entire document | • | | | | | | |
| | o, pages so a co, see came accument. | | | | | | | |
| X | US 6,008,243 A (BENDER et al.) 28 D | ecember 1999(28.12.99), see | 1-7, 15-20 | | | | | |
| | entire document. | | | | | | | |
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| Further documents are listed in the continuation of Box C. See patent family annex. | | | | | | | | |
| * Special categories of cited documents: "A" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention | | | | | | | | |
| "A" do to | invention | | | | | | | |
| | rlier document published on or after the international filing date cument which may throw doubts on priority claim(s) or which is | "X" document of particular relevance; the considered novel or cannot be considered when the document is taken alone | | | | | | |
| cit | e claimed invention cannot be | | | | | | | |
| *O* do | ecial reason (as specified) cument referring to an oral disclosure, use, exhibition or other cans | step when the document is a documents, such combination | | | | | | |
| ¹P¹ do | document published prior to the international filing date but later than "&" document member of the same patent family the priority date claimed | | | | | | | |
| Date of the actual completion of the international search Date of mailing of the international search report | | | | | | | | |
| 12 ЛЛГА | 2001 | 30 JUL 2007' | | | | | | |
| Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Authorized offices Authorized offices Authorized offices | | | | | | | | |
| Box PCT | oner of Patents and Trademarks | AMY J. HARTTER | TUN | | | | | |
| Washington, D.C. 20231 Facsimile No. (703) 305-3230 | | Telephone No. (703) 308-0196 | | | | | | |

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US01/05150

| Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet) | | | | | | |
|--|--|--|--|--|--|--|
| This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons: | | | | | | |
| 1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: | | | | | | |
| 2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: | | | | | | |
| 3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). | | | | | | |
| Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet) | | | | | | |
| This International Searching Authority found multiple inventions in this international application, as follows: | | | | | | |
| Please See Extra Sheet. | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| | | | | | | |
| 1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. | | | | | | |
| 2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. | | | | | | |
| 3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: | | | | | | |
| | | | | | | |
| 4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: | | | | | | |
| Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees. | | | | | | |

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/05150

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for as inventions to searched the appropriate search fees must be paid.

Group I which consists of claims 1-7 is distinct as it addresses itself to the solution complex of the mixture of MMP-13 and the defined "Compound A." The solution is clearly distinct and different from the crystal complex, active site and methods that are claimed in succeeding groups and according claims.

Group II consists of claims 8-14. These claims pertain to the actual product of the crystal complexion its entirety. Thus it is distinct from Groups I and Groups 3-4. The group claims the whole crystal known as "Compound A" and the crystal is not in any other type of alternate environment or with any additional accountrements.

Group III encompasses the claims of 15-20. These claims consist of the active site of the molecule of MMP-13. This chemical is a portion of the solution claimed in the first group and thus separate and distinct from the solution of Group I or the separate entity of "Compound A" that is claimed in Group 2. Thus these Groups are separate.

Group IV consists of claims 21-32 which claim a method of identifying an inhibitor or activator of the MMP-13 compound. The method that is embodied within this Group is clearly different from the proceeding groups. Firstly the claims within Group 4 are directed toward a method of accomplishing the task of identifying different entities and not a product itself. Secondly its actions are addressed to entities outside the compound itself and not limited to "Compound A" of the MMP-13. Based on the aforementioned reasons and the distinct nature of the claims defined in each of the groups, the instant application has a lack of unity due to each group having a different Special Technical Peature a summarized above for each group.

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